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FEDERAL REPUBLIC OF GERMANY [Eagle crest]



Priority Certificate for the filing of a Patent Application

File Reference:

199 26 068.0

Filing date:

8 June 1999

Applicant/Proprietor: Professor Dr. Arne S k e r r a, Freising/DE

Title:

Muteins of the bilin-binding protein

IPC:

C 07 K, C 12 N and C 12 Q

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Patent Specification DE 199 26 068 C 1 11

Int. Cl.⁷: C 07 K 14/435 C 12 N 15/62 C 12 Q 1/68

[crest]

GERMAN PATENT OFFICE

21 File reference 199 26 068.0-41

22 Date of application 8.6.1999

43 Date laid open

46 Date of publication of the grant of the patent: 11.1.2001

Opposition can be notified within 3 months from the publication of grant

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56 Printed publications taken into consideration for assessing patentability:

Eur. J. Biochem. 219, S. 855-863, 1994;

Muteine des Bilin-Bindungsproteins

Die Erfindung bezieht sich auf Muteine des Bilin-Bindungsproteins mit Bindungsfähigkeit für Digoxigenin sowie Fusionsproteine solcher Muteine, Verfahren zur Herstellung derartiger Muteine und ihrer Fusionsproteine sowie deren Verwendung zum Nachweis oder zur Bindung von mit Digoxigenin markierten Biomolekülen. Insbesondere betrifft die Erfindung ein Polypeptid, ausgewählt aus Muteinen des Bilin-Bindungsproteins, welches dedurch gekennzeichnet ist, daß es (a) Digoxigenin oder Konjuga te des Digoxigenins zu binden vermag, (b) Ouabain, Testosteron und 4-Aminofluorescein nicht bindet und (c) an mindestens einer der Sequenzpositionen 28, 31, 34, 35, 36, 37, 58, 60, 69, 88, 90, 95, 97, 114, 116, 125 und 127 des Bilin-Bindungsproteins eine Aminosauresubstitution aufweist. Aufgrund ihres einfachen molekularen Aufbaus weisen die erfindungsgemäßen Muteine bei Herstellung und Verwendung Vorteile im Vergleich zu Antikörpern gegen die Digoxigeningruppe auf.

BUNDESDRUCKEREI 11.00 002 162/146/7A

DE 199 26 068 C 1

DE 199 26 068 C1



Description

The present invention relates to muteins of the bilinbinding protein which are capable of digoxigenin and to fusion proteins of such muteins, to methods for preparing muteins of this kind and their also to the use thereof fusion proteins and binding biomolecules labeled 10 detecting or with digoxigenin.

In molecular biology, the digoxigenin group is these a very common instrument for nonradioactive days nucleic acids, proteins detection οf and biomolecules. For this purpose, the biomolecule is, mostly covalently, modified with a reactive digoxigenin derivative, thus allowing subsequent detection of the using an antibody directed against molecule digoxigenin group or a conjugate of an appropriate antibody fragment and a reporter enzyme, according to generally used methods in biochemistry.

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The skilled worker knows quite a number of reactive 25 digoxigenin derivatives which are partially commercially available. For example, digoxigenin-3-0methylcarbonyl-ε-aminocaproic acid N-hydroxysuccinimide (DIG-NHS), digoxigenin-3-O-succinyl-ε-aminocaproic acid N-hydroxysuccinimide ester and 3-amino-3deoxydigoxigenin-hemisuccinimide succinimidyl ester are 30 suitable for covalent coupling to proteins, particular to the amino groups of exposed lysine side chains. Using 3-iodoacetylamino-3-deoxydigoxigenin possible to label especially thiol groups in proteins or in other biomolecules selectively with the 35 digoxigenin group. It is possible to couple synthetic oligodeoxynucleotides to the same reactive digoxigenin derivatives, as long as they have been provided with suitable free amino or thiol groups during synthesis.

addition, cis-platinum complexes of digoxigenin derivatives (DIG Chem-Link reagent) or digoxigenin derivatives containing carbodiimide groups (disclosed in the European patent specification EP 0 806 431 A2) suitable for direct labeling of nucleic acids. Alternatively, it is possible in the deoxyribonucleic acids to label said deoxyribonucleic acids during a matrix-dependent enzymic synthesis with aid of a DNA polymerase and a deoxynucleotide triphosphate coupled to the digoxigenin group, example digoxigenin-11-dUTP, digoxigenin-11-ddUTP digoxigenin-16-dATP. Analogously, digoxigenin-11-UTP is suitable for incorporation into enzymically synthesized RNA. Moreover, it is possible to label oligodeoxynucleotides with the digoxiqenin directly in the automated DNA synthesis by suitable activated building blocks, for example nucleotides". Digoxigenin "virtual group-coupled nucleic acids of this kind are suitable as for nonradioactive gene probes detection of complementary nucleotide sequences by hybridization, for example in Northern or Southern blots (disclosed in the European patent specification EP 0 324 474 A1).

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25 Digoxigenin group-labeled proteins or glycoproteins are particularly useful for determining, for example, relevant antigens or antibodies directed thereagainst in immunochemical assay methods such as ELISA (enzymelinked immunosorbent assay). The biomolecule conjugated 30 with the digoxigenin group is actually detected using an anti-digoxigenin antibody, normally in the form of a conjugate of the Fab fragment of said antibody with a for example, alkaline suitable enzyme, such as, phosphatase or horseradish peroxidase, as label. 35 enzymic activity then serves for quantification chromogenic, fluorogenic catalysis of а or chemiluminescent reaction. Various antibodies against the digoxigenin group are known (Mudgett-Hunter et al.,

J. Immunol. 129 (1982), 1165-1172; Jeffrey et al., J. Mol. Biol. 248 (1995), 344-360).

of antibodies, however, has The use of monoclonal disadvantages. Thus, preparation antibodies in hybridoma cell cultures is complicated, and proteolysis to give the Fab fragment and also production of conjugates with reporter enzymes requires additional difficult process steps. But even production of antibodies by genetic engineering is not 10 simple, and the main reason for this is that antibodies as antigen-binding fragments thereof are composed of two different polypeptide chains in a genetic complicated manner. For structurally manipulation of antibodies it is therefore necessary to 15 handle two genes simultaneously. Moreover, the yield of correctly folded antibody fragments produced by genetic engineering is often low. As is known to the skilled worker, this is even more so when recombinant fusion proteins are to be prepared from Fab fragments of 20 antibodies and enzymes.

It was therefore the object of the invention to develop alternative polypeptide reagents for detection of the digoxigenin group, which can be produced in a simple manner.

In an evolutive research approach, it has surprisingly been found now that muteins of the bilin-binding protein which is structurally based on a single polypeptide chain (Schmidt and Skerra, Eur. J. Biochem. 219 (1994), 855-863) are suitable for detecting the digoxigenin group by high-affinity binding, with digoxigenin recognition being astoundingly selective compared with other steroids.

The present invention thus relates to a polypeptide, selected from muteins of the bilin-binding protein, which is characterized in that it

- (a) is able to bind digoxigenin or digoxigenin conjugates,
- (b) does not bind ouabain, testosterone and 4-aminofluorescein and
- (c) has an amino acid substitution at at least one of the sequence positions 28, 31, 34, 35, 36, 37, 58, 60, 69, 88, 90, 95, 97, 114, 116, 125 and 127 of the bilin-binding protein.

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- Outside the region of amino acid positions 28, 31, 34, 10 35, 36, 37, 58, 60, 69, 88, 90, 95, 97, 114, 116, 125 and 127 the muteins of the present invention may correspond to the amino acid sequence of the bilinbinding protein from Pieris brassicae. On the other 15 hand, the amino acid sequence of the polypeptides of the invention may have differences to the bilin-binding protein also besides said positions. Bilin-binding variants of this kind protein sequence comprise naturally occurring and also artificially generated the deviations mean substitutions, 20 variants. and insertions, deletions of amino acid residues and also N- and/or C-terminal additions.
- For example, the inventive muteins of the bilin-binding 25 protein may have amino acid substitutions which prevent oligomerization of the bilin-binding protein, such as the Asn(1)->Asp substitution, or suppress proteolytic cleavage within the polypeptide chain, which may occur during production in E. coli, such as, for example, the 30 Lys(87)->Ser substitution. Furthermore, it is possible mutations Asn(21)->Gln introduce the Lys(135)->Met into the nucleic acids coding for the muteins of the bilin-binding protein, in order to facilitate, for example, cloning of a gene segment via two new BstXI restriction cleavage sites at these 35 positions. Likewise, the present invention relates to the specific introduction of amino acid substitutions within or outside the said positions, in order to generally improve particular properties of the mutein

of the invention, for example its folding stability or folding efficiency or its resistance to proteases.

The ability of the polypeptides of the invention to bind digoxigenin or digoxigenin conjugates be by common methods, for example determined fluorescence titration, titration calorimetry, surface plasmon resonance measurements or blotting methods, for example Western blotting, Southern blotting or Northern blotting. Blotting methods may be used in order to 10 transfer conjugates of digoxigenin with proteins nucleic acids to a membrane and then detect said conjugates using one of the muteins of the invention, a conjugate of this mutein or a fusion protein of this mutein. 15

A quantitative parameter for binding affinity is provided by established thermodynamic parameters such as, for example, the affinity constant or dissociation constant for the complex of mutein and bound ligand, for example digoxigenin. However, it is also possible to determine the binding ability qualitatively, for example based on the intensity of a binding signal due to a chromogenic reaction or of a colored precipitate which is obtained with the aid of one of said blotting methods.

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Preferred muteins of the invention are obtainable in a two-stage evolutive process. Random mutagenesis of the bilin-binding protein and repeated selection of muteins with digoxigenin group affinity from this library, using free digoxigenin for competitive concentration, provides muteins of the bilin-binding protein which recognize the digoxigenin group, but the affinity is still comparatively low. Renewed mutagenesis of such a mutein at amino acid positions 28, 31, 34, 35, 36 and 37, now followed by a repeated concentration by formation of a complex with the digoxigenin group and by subsequent dissociation of the complex formed in

acidic medium, then results in obtaining muteins having substantially higher affinity for the digoxigenin group.

5 Surprisingly, it has now been found that the affinity constant between such polypeptides of the invention and digoxigenin is at least 10⁷ M⁻¹. This means in other words that the dissociation constant of the complex of the polypeptide of the invention and digoxigenin is 10 100 nM or less. Individual examples even show dissociation constants of 35 nM or less, as illustrated in the examples.

digoxigenin, the inventive muteins of also bind · 15 bilin-binding protein can digoxigenin derivatives as ligands, for example digoxin, digitoxin or digitoxigenin. Furthermore, the inventive muteins of the bilin-binding protein may bind conjugates of said chemical compounds, i.e. nucleic acids, polypeptides, 20 carbohydrates, other natural or synthetic biomolecules, macromolecules or low molecular weight compounds which are covalently linked or linked via a metal complex to digoxin, digitoxin or digoxigenin, digitoxigenin. Preference is given to using for the preparation of 25 the reactive derivatives such conjugates of digoxigenin, digoxin, digitoxin or digitoxigenin, which are known to the skilled worker and are stated, for example, further above.

30 Preferred muteins of the invention, which were obtained by the two-stage process described, show, compared with the affinity for digoxigenin, an even higher affinity for digitoxin or digitoxigenin whose steroid system differs from that of digoxigenin only by the absence of a hydroxyl group. Surprisingly, these muteins show distinctive specificity with respect to the digoxigenin or digitoxigenin group, and this is shown by the fact that other steroids or steroid groups such as ouabain or testosterone are bound with much less affinity, if

at all. Fluorescein derivatives such as 4-aminofluorescein, too, are evidently not bound. This means that ouabain, testosterone or 4-aminofluorescein in each case have a dissociation constant of at least 10 μ M, preferably at least 100 μ M for the inventive muteins of the bilin-binding protein.

This property of specificity distinguishes said muteins considerably from other muteins of the bilin-binding 10 protein and also from antibodies directed against the digoxigenin group, such as, for example, antibody 26-10 (Chen et al., Protein Eng. 12 (1999), 349-356) which binds ouabain with substantial affinity, and gives the inventive muteins of the bilin-binding protein a advantage. Ιt is surprising 15 particular particularly the additional amino acid substitutions at 35, 36 and 37 lead to the positions 28, 31, 34, preferred muteins of the bilin-binding protein. Preference is therefore given to those muteins which 20 least one or all of the amino carry at substitutions Glu(28)->Gln, Lys(31)->Ala, Asn(34)->Asp, $Ser(35) \rightarrow His$, $Val(36) \rightarrow Ile$ and $Glu(37) \rightarrow Thr$.

Particularly preferred muteins of the invention carry, when compared to the biling-binding protein, at least 25 the amino acid substitutions selected from $Glu(28) \rightarrow Gln$, Lys(31) -> Ala, Asn(34) -> Asp, Ser(35) -> His, Val(36) - Sile, Glu(37) - Shr, Asn(58) - Arg, His(60) - Ser, Ile(69) - Ser, Leu(88) - Tyr, Tyr(90) - Ile, Lys(95) - Gln, 30 Asn(97) - Sly, Tyr(114) - Phe, Lys(116) - Ser, Gln(125) - Pheand Phe(127)->Leu. The representation indicates in each case first the amino acid in the natural bilin-binding protein (SWISS-PROT accession code P09464) together with the sequence position for the mature polypeptide in brackets, and 35 corresponding amino acid in a mutein of invention is stated after the arrow. Very particularly preferred muteins according to this invention carry all of the amino acid substitutions mentioned.

Surprisingly, bilin-binding protein position 93 is unchanged in the muteins of the invention, although this amino acid, too, had been affected by the mutagenesis for preparing the random library. Preferred muteins of the bilin-binding protein therefore carry the amino acid Val at said position.

It is an advantage for particular detection methods to use the muteins of the bilin-binding protein of the present invention in a labeled form. Accordingly, this 10 invention further relates to a polypeptide of the invention, which is characterized in that it carries at least one label. Suitable labeling groups are known to skilled worker and include enzymic label, radioacctive label, fluorescent label, chromophoric 15 label, (bio) luminescent label or a label containing haptens, biotin, metal complexes, metals or colloidal Very generally, labeling is possible substances or enzymes which generate a determinable substance in a chemical or enzymatic reaction. In this 20 connection it is possible to couple all known labels for antibodies to the muteins of the invention, too.

A possibility which is particularly advantageous for practical application is to use the inventive muteins of the bilin-binding protein in the form of fusion proteins. Techniques for preparing such fusion proteins by means of genetic engineering methods are known to the skilled worker. Suitable fusion partners for the muteins of the invention would be enzymes and other polypeptides, proteins or protein domains. Such fusions would be suitable for providing the mutein of the bilin-binding protein with additional properties such as, for example, enzymic activity or affinity for other molecules such as proteins, macromolecules or low molecular weight ligands.

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Examples of possible fusions are those with enzymes which catalyze chromogenic or fluorogenic reactions or

release of cytotoxic may be used for the agents. Further examples for fusion partners which may advantageous in practice are binding domains such as albumin-binding domain or the immunoglobulinbinding domain of protein G or protein A, antibody fragments, oligomerization domains, toxins or other binding proteins and functional parts thereof and also affinity peptides such as, for example, Strep tag or Strep tag II (Schmidt et al., J. Mol. Biol. 255 (1996), 10 753-766). Suitable fusion partners are also proteins chromogenic or fluorogenic particular having properties, such as, for example, green fluorescent protein. Another suitable fusion partner would be coat protein III of a filamentous bacteriophage such as M13, fl or fd, or a fragment of said coat protein. 15

Very generally, the term fusion protein is intended here to mean also those inventive muteins of the bilinbinding protein, which are equipped with a sequence. Signal sequences on the N terminus of the polypeptide of the invention may serve the purpose of directing said polypeptide during biosynthesis into a particular cell compartment, for example the E. coli periplasm or the lumen of the endoplasmic reticulum of a eukaryotic cell, or into the medium surrounding the cell. The signal sequence is typically cleaved off in the process by a signal peptidase. In addition, it is signal possible use other ortargeting to sequences which need not necessarily be located on the N terminus of the polypeptide and which make possible to locate said polypeptide in specific cell compartments. A preferred signal sequence for secretion into the E. coli periplasm is the ompA signal sequence. A large number of further signal sequences and also targeting sequences are known in the prior art.

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An advantage of the inventive muteins of the bilinbinding protein is the suitability of both their N terminus and their C terminus for preparing fusion

In contrast to antibodies in which proteins. the light and heavy N terminus of both the immunoglobulin chain is in spatial proximity to antigen binding site, it is possible to use in invention of polypeptides of the both ends the polypeptide chain for the preparation proteins, without adversely affecting ligand binding.

The invention therefore also relates to fusion proteins of muteins of the bilin-binding protein, in which an 10 enzyme, another protein or a protein domain, a signal sequence and/or an affinity peptide is fused to the amino terminus of the polypeptide in an operable manner. The invention yet further relates to fusion 15 proteins of bilin-binding protein muteins or of fusion proteins having the amino terminus of biling-binding porotein muteins, in which an enzyme, another protein or a protein domain, a targeting sequence and/or an affinity peptide is fused to the carboxy terminus of the polypeptide in an operable manner. 20

A preferred enzyme for constructing the fusion proteins invention is bacterial alkaline phosphatase of the (Sowadski et al., J. Mol. Biol. 186 (1985) 417-433) which may be attached either at the N terminus or at the C terminus of a mutein of the bilin-binding protein. In addition, such a fusion protein may carry a signal sequence such as, for example, OmPA or PhoA, which effect secretion of said fusion protein into the E. coli periplasm where the disulfide bonds may form efficiently in the polypeptide chain. Furthermore, may be equipped with an affinity peptide such as, for example, Strep tag II, which allows easy purification of said fusion protein. The specific fusion proteins of invention are described in the examples. the advantage of a fusion protein of this kind is ability to catalyze directly a chromogenic, fluorogenic chemiluminescent detection reaction, which or

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simplifies its use for detection of the digoxigenin group.

Another advantage of using alkaline phosphatase for constructing fusion proteins of the invention is the fact that this enzyme forms a stable homodimer and, consequently, confers the property of bivalence on the bilin-binding protein mutein as part of the fusion protein. In this way, binding of the digoxigenin group may result in avidity effect which increases detection 10 sensitivity. Such an avidity effect can be expected in particular if the digoxigenin-labeled molecule adsorbed to a solid phase, is present in oligomeric or membrane-bound form or is conjugated with a plurality 15 of digoxigenin groups. Analogously, other homodimeric enzymes are suitable for preparing bivalent fusion proteins containing the inventive muteins of the bilinbinding protein.

20 Apart from bacterial alkaline phosphatase, it is also possible to use phosphatases from eukaryotic organisms, such as, for example, calf intestine phosphatase (CIP), for preparing fusion proteins of the invention. Said phosphatases are frequently distinguished by higher 25 enzymatic activity (Murphy Kantrowitz, and Microbiol. 12 (1994), 351-357), which may result higher detection sensitivity. It is also possible to use mutants of bacterial alkaline phosphatase, which have improved catalytic activity (Mandecki et al., Protein Eng. 4 (1991), 801-804), for constructing 30 fusion proteins of the invention. Other enzymes known to the skilled worker which catalyze chromogenic, fluorogenic or chemiluminescent reactions, such as, for example, β -galactosidase or horseradish peroxidase, are also suitable for preparing fusion proteins of the 35 invention. Moreover, all these enzymes may likewise be employed for labeling muteins of the bilin-binding protein by conjugating them, for example by using common coupling reagents, with the mutein obtained separately or a fusion protein of the mutein.

In another aspect, the present invention relates to a nucleic acid which comprises a sequence coding for a mutein or a fusion protein of a mutein of the bilinbinding protein. This nucleic acid may be part of a vector which contains operatively functional areas for expressing the nucleic acid. A large number of suitable vectors is known from the prior art and 10 described in detail here. Operatively functional areas are those elements which allow, assist, facilitate increase and/or transciption and/or subsequent processing of an mRNA. Examples of elements of this 15 kind include promoters, enhancers, transcription initiation sites and transcription termination sites, translation initiation sites, polyadenylation signals, etc.

20 The nucleic acid of the invention or its surrounding areas may be such that biosynthesis of the polypeptide takes place in the cytosol, the polypeptide sequence preceded, where appropriate, by starting a methionine. In a preferred embodiment, however, 25 N-terminal signal sequence is used, in particular the OmpA or PhoA signal sequence, in order to direct the polypeptide of the invention into the E. coli periplasm where the signal sequence is cleaved off by the signal peptidase and the polypeptide chain is able to fold 30 with oxidative formation of the disulfide bonds. Eukaryotic signal sequences may be used in order to secrete the polypeptide of the invention eukaryotic host organism. In principle, both prokaryotic, preferably E. coli, and eukaryotic cells 35 such as, for example, yeasts are considered expression of the nucleic acid of the invention.

In yet another aspect, the present invention relates to a method for preparing an inventive mutein or fusion protein of a mutein of the bilin-binding protein, which method is characterized in that the nucleic acid coding for the mutein or the fusion protein of a mutein of the bilin-binding protein is expressed in a bacterial or eukaryotic host cell and the polypeptide is obtained from the cell or the culture supernatant. For this purpose, normally a suitable host cell is first transformed with a vector which comprises a nucleic acid coding for a polypeptide of the invention. The host cell is then cultured under conditions under which the polypeptide is biosynthesized, and the polypeptide of the invention is obtained.

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With respect to the preparation method, it must be taken into account that the inventive muteins of the 15 bilin-binding protein have two structural disulfide bonds and that additional disulfide bonds may present in corresponding fusion proteins. The formation of said disulfide bonds, which takes place during protein folding, is normally ensured if the polypeptide 20 of the invention is directed with the aid of a suitable signal sequence into a cell compartment containing an oxidizing thiol/disulfide redox medium, for into the bacterial periplasm or the lumen of the 25 endoplasmic reticulum of a eukaryotic cell. In this connection, the polypeptide of the invention can be liberated by cell fractionation or obtained from the culture supernatant. It is possible, where appropriate, to increase the folding efficiency by overproducing protein disulfide isomerases, for example E. coli DsbC 30 protein, or auxiliary folding proteins.

On the other hand, it is possible to produce a polypeptide of the invention in the cytosol of a host cell, preferably *E. coli*. The said polypeptide may then be obtained, for example, in the form of inclusion bodies and then be renatured *in vitro*. Depending on the intended use, the protein can be purified by means of various methods known to the skilled worker. A suitable

method for purifying the inventive muteins of the bilin-binding protein is, for example, affinity chromatography using a column material which carries digoxigenin groups. In order to purify fusion proteins of the muteins of the bilin-binding protein, it possible to utilize the affinity properties of the fusion protein, which are known from the prior art, for example those of the Strep tag or the Strep tag II (Schmidt and Skerra, J. Chromatogr. A 676 (1994), 337-345; Voss and Skerra, Protein Eng. 10 (1997), 975-982), 10 those of the albumin binding domain (Nygren et al., J. Mol. Recogn. 1 (1988), 69-74) or those of alkaline phosphatase (McCafferty et al., Protein Eng. 4 (1991) 955-961). The fact that the muteins of the bilinbinding protein consist only of a single polypeptide 15 chain is advantageous for the methods for preparing the polypeptides of the invention, since no care needs to taken either of the need for synthesizing a plurality of various polypeptide chains within a cell of different polypeptide chains 20 simultaneously or associating with one another in a functional manner.

practical application possibilities for The of bilin-binding inventive muteins the protein essentially correspond to those for conventional antibodies or antibody fragments with binding affinity the digoxigenin. Accordingly, invention relates to the use of a mutein of the invention or of a fusion protein of a mutein of the bilin-binding protein in a method for detecting, determining, immobilizing or removing digoxigenin or conjugates of digoxigenin with proteins, nucleic acids, carbohydrates, biological or synthetic macromolecules or low molecular weight chemical compounds.

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The inventive muteins of the bilin-binding protein or their fusion proteins can be used in detection methods essentially in a manner analogous to corresponding detection methods known for anti-digoxigenin antibodies and also fragments and so-called conjugates thereof. In a further aspect, the present invention therefore relates to a method for detecting the digoxigenin group, in which method a mutein of the bilin-binding protein or a fusion protein of a mutein of the bilin-binding protein is contacted with digoxigenin or with digoxigenin conjugates under conditions suitable for effecting binding of the mutein to the digoxigenin group and the mutein or the fusion protein of the mutein is determined.

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For this purpose, the mutein may be labeled directly, for example by covalent coupling. It is, however, also possible to use indirect labeling, for example by means of labeled antibodies against the bilin-binding protein muteins thereof or against domains proteins of these muteins. The use of inventive fusion proteins containing an enzyme, for example alkaline phosphatase, instead of a labeled mutein of the bilinbinding protein is particularly advantageous. In this case, it is possible to design the determination method with a particularly small number of process steps, it being possible to utilize directly, for example, the ability of the enzyme as part of the fusion protein to catalyze a chromogenic, fluorogenic or luminescent detection reaction. Here, the fact that such fusion readily available is a particular proteins are advantage compared with corresponding fusion proteins antibodies. Utilization of conventional abovedescribed avidity effect in the case oligomeric fusion protein is another advantage in such a method.

It is possible to carry out a method for determining the digoxigenin group, for example, qualitatively for detecting nucleic acids conjugated with the digoxigenin group in Southern or Northern blots or proteins conjugated with the digoxigenin group in Western blots. A determination method may also be carried out

quantitatively for detecting proteins conjugated with the digoxigenin group in an ELISA. In addition, a determination method of the invention is also suitable for indirect detection of proteins not conjugated with digoxigenin or of other molecules by using a binding is directed against the protein protein which molecule, for example an antibody or its fragment, and conjugated with the digoxigenin is Indirect detection of the nucleic acids not conjugated with digoxigenin is also possible by using a gene probe which hybridizes with said nucleic acid and which is conjugated with the digoxigenin group. An application in medical diagnostics or therapy results in addition from determination of digoxigenin, digoxin, digitoxin or digitoxigenin, without these ligands having to be conjugated with another molecule.

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The muteins of the invention or fusion proteins thereof may also be used for immobilizing a molecule conjugated with the digoxigenin group. This immobilization is preferably carried out on solid phases coated with the muteins or their fusion proteins, such as, for example, microtiter plates, immunosticks, microbeads made of organic, inorganic or paramagnetic materials, or sensor surfaces.

Correspondingly, it is likewise possible to use the muteins of the invention or fusion proteins thereof for removing digoxigenin, digoxin, digitoxin digitoxigenin, or a molecule conjugated with one of these compounds. In this case, in addition to the solid phases mentioned, column materials are also considered for coating with the muteins or their fusion proteins. Preferably, said coating is carried out on suitable column materials by coupling by means of chemically reactive groups. Column materials coated in this way may be used for removing substances conjugated with digoxigenin groups and also, where appropriate,

complexes of such substances with other molecules from a solution.

Thus, it is possible, for example, to remove antigens from a solution by adding antibodies to the solution, which are directed against the antigens and conjugated digoxigenin group, and with the contacting resulting solution with said column material under conditions under which a complex between the digoxigenin groups and an inventive mutein of 10 bilin-binding protein or its fusion protein is formed. Following such a removal, it is also possible, where appropriate, to elute the substance conjugated with digoxigenin. This elution may be carried out competition with digoxin, digoxigenin, digitoxin or 15 digitoxigenin and also, for example, by lowering or increasing the pH of the solution. In a competitive elution it is possible to utilize in an advantageous manner the higher binding affinity of the muteins of 20 the invention for digitoxigenin or digitoxin compared with the digoxigenin group. In this way it is possible isolate or purify a substance conjugated with digoxigenin.

- 25 The invention is further illustrated by the following examples and attached drawings, in which:
- Fig. 1 represents in each case a fluorescent titration of the Strep tag II-fused mutein DigA16 with the ligands digoxigenin, digitoxigenin and ouabain;
- Fig. 2 depicts diagrammatically the expression vectors pBBP27 (A) and pBBP29 (B) for preparing fusion proteins of mutein DigA16 with alkaline phosphatase;
 - Fig. 3 demonstrates quantitative detection of biomolecules conjugated with digoxigenin

groups by fusion proteins of mutein DigA16 with alkaline phosphatase in an ELISA;

- Fig. 4 shows qualitative detection of biomolecules conjugated with digoxigenin groups by fusion proteins of mutein DigA16 with alkaline phosphatase on a Western blot.
- Fig. 1 shows the graphic representation of results from Example 3 in which different concentrations of the 10 steroids digoxigenin (squares), digitoxigenin (circles) and ouabain (rhomboids) were added to a 1 µM solution of mutein DigA16. The particular protein fluorescence intensities were measured at an excitation wavelength of 295 nm and an emission wavelength of 345 nm and 15 function of the actual total steroid plotted as a in the particular reaction mixture. concentration Finally, the data points were fitted to a regression curve by means of nonlinear regression.

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Fig. 2 shows a drawing of the expression vectors pBBP27 (A) and pBBP29 (B). pBBP27 codes for a fusion protein of bacterial alkaline phosphatase with its own signal sequence, a peptide linker having the sequence Pro-Pro-Ser-Ala, the mutein DigA16 and also the Strep tag II 25 affinity tag. The corresponding structural gene followed by the dsbC structural gene (including its ribosomal binding site) from E. coli (Zapun et al., Biochemistry 34 (1995), 5075-5089) as second cistron. 30 The artificial operon formed in this way is under joint of the tetracyclin transcriptional control promoter/operator (tet^{p/o}) and ends at the lipoprotein transcription terminator (t_{1pp}) . Further vector elements are the origin of replication (ori), the intergenic 35 region of filamentous bacteriophage f1 (f1-IG), the ampicillin resistance gene (bla) coding for β -lactamase and the tetracyclin repressor gene (tetR). pBBP29 codes for a fusion protein of the OmpA signal sequence, the mutein DigA16, the Strep tag II affinity tag, a peptide

linker consisting of five glycine residues, and bacterial alkaline phosphatase without its N-terminal amino acid arginine. The vector elements outside this region are identical to vector pBBP27.

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Fig. 3 shows a graphic representation of the data from Example 4 in which digoxigenin groups were detected quantitatively with the aid of mutein DigAl6 fusion proteins as gene products of vectors pBBP27 (closed symbols). and pBBP29 (open Here, symbols) digoxigenin groups were coupled on the one hand to bovine serum albumin (BSA, squares) or, on the other to albumin (ovalbumin, triangles). hand, egg control data shown are those obtained when using underivatized bovine serum albumin and the fusion protein encoded by pBBP27 (open circles). The enzymic activity corresponding to the particular bound fusion protein was monitored spectrophotometrically at 405 nm on the basis of p-nitrophenyl phosphate hydrolysis. Curve fitting was carried out by nonlinear regression with the aid of the Kaleidagraph computer program (Abelbeck Software) by means of the equation

 $[P \cdot L] = [L]_t [P]_t / (K_d + [P]_t)$.

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[P]t corresponds to the total fusion protein concentration used in the particular microtiter plate well. [P•L] is determined on the basis of the enzymic phosphatase. The total activity of alkaline concentration of digoxigenin groups 30 [L]_t, constant well concentration series, per and within a dissociation constant K_d were fitted as parameters by nonlinear regression.

Fig. 4 shows the result of a Western blot experiment from Example 4 for qualitative detection of biomolecules conjugated with digoxigenin groups by means of the mutein DigA16 fusion proteins encoded by pBBP27 (lanes 1 and 2) and pBBP29 (lanes 3 and 4). For

comparison, a 15% strength SDS polyacrylamide gel of the biomolecules, stained with Coomassie Brilliant Blue, is likewise shown (lanes 5 and 6). Here, a mixture of 0.5 µg of underivatized BSA, underivatized ovalbumin and underivatized RNaseA was fractionated in each case in lanes 1, 3 and 5. A mixture of 0.5 µg of BSA coupled to digoxigenin groups, ovalbumin coupled to digoxigenin groups and RNaseA coupled to digoxigenin groups was fractionated in each case in lanes 2, 4 and 6.

Examples

Unless stated otherwise, the genetic engineering methods familiar to the skilled worker, as described, for example, in Sambrook et al. (Molecular Cloning. A Laboratory Manual (1989), Cold Spring Harbor Press) were used.

20 Example 1

Preparation of a library for muteins of the bilinbinding protein, phagemid presentation and selection of a mutein with binding affinity for digoxigenin

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A library for muteins of the bilin-binding protein was the amino acid prepared by subjecting positions of said bilin-binding protein, 34, 35, 37, 58, 60, 69, 88, 90, 93, 95, 97, 114, 116, 125 and 127, to a concerted mutagenesis in multiple steps with the aid of the polymerase chain reaction (PCR). The PCR reactions were initially carried out in two separate amplification steps in a volume of in each case 50 µl, and 10 ng of pBBP20 phasmid DNA (SEQ ID NO: 1) matrix and in each case 25 pmol of two primers (SEQ ID NO. 2 and SEQ ID NO. 3 in one mixture and SEQ ID NO. 4 and SEQ ID NO. 5 in a second mixture) which had been synthesized according to the generally phosphoramidite method were used.

Furthermore, the reaction mixture contained 5 µl of 10xTaq buffer (100 mM Tris/HCl pH 9.0, 500 mM KCl, 1% v/v Triton X-100), 3 μ l of 25 mM MgCl₂ and 4 μ l of dNTP mix (2.5 mM dATP, dCTP, dGTP, dTTP). After filling up with water, the mixture was overlaid with mineral oil and heated to 94°C in a programmable thermostating block for 2 min. Then 2.5 u of Taq DNA polymerase (5 $u/\mu l$, Promega) were added and 20 temperature cycles of 1 min at 94°C, 1 min at 60°C and 1.5 min at 72°C were carried out, followed by an incubation at 60°C for 5 min. The desired amplification products were isolated via preparative agarose gel electrophoresis from low melting point agarose Jetsorb (Gibco BRL), using the according DNA extraction kit (Genomed) to the manufacturer's instructions.

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A relevant section of the pBBP20 nucleic acid sequence is shown together with the encoded amino acid sequence as SEQ ID NO. 1 in the sequence listing. The section starts with a hexanucleotide sequence which was obtained by ligating an XbaI overhang with an SpeI overhang complementary thereto and ends with the HindIII cleavage site. The vector elements outside this region are identical to vector pASK75 whoses complete nucleotide sequence is stated in the publication DE 44 17 598 A1.

The subsequent amplification step was carried out in a 100 µl mixture, and in each case approx. 6 ng of the two isolated fragments as matrix, 50 pmol of each of the two primers SEQ ID NO. 6 and SEQ ID NO. 7 and also 1 pmol of oligodeoxynucleotide SEQ ID NO. 8 were used. The remaining components of the PCR mixture were added in twice the amount, as in the preceding amplification steps. The PCR was carried out in 20 temperature cycles of 1 min at 94°C, 1 min at 55°C, and 1.5 min at 72°C, followed by a final incubation at 60°C for 5 min. The fragment obtained was again isolated by preparative agarose gel electrophoresis.

Said fragment which represented the mutein library in the form of a mixture of nucleic acids was cloned by cutting it first with the restriction enzyme BstXI (New Biolabs) according to the manufacturer's England instructions. The nucleic acid fragment obtained (335 bp) was purified again by base pairs, means preparative agarose gel electrophoresis. Analogously, pBBP20 vector DNA was cut with BstXI and the larger of the two fragments (3971 bp) was isolated.

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For ligation, 0.93 µg (4.2 pmol) of the PCR fragment and 11 µg (4.2 pmol) of the vector fragment were incubated in the presence of 102 Weiss units of T4 DNA ligase (New England Biolabs) in a total volume of 500 μl (50 mM Tris/HCl pH 7.8, 10 mM MgCl, 10 mM DTT, 1 mM ATP, 50 μ g/ml BSA) at 16°C for two days. The DNA was then precipitated by adding 10 µg of yeast tRNA (Boehringer Mannheim), 25 µl of 5 M ammonium acetate and 100 μ l of ethanol to in each case 24 μ l of the ligation mixture. Incubation at -20°C for 3 days was followed by centrifugation (25 min, 16000 g, 4°C). The precipitate was washed in each case with 200 µl (70% v/v, −20°C) and dried under ethanol pressure. Finally, the DNA was taken up in 43.6 μ l of TE/10 (1 mM Tris/HCl pH 8.0, 0.1 mM EDTA). The DNA concentration of the solution obtained was estimated by analytical agarose gel electrophoresis on the basis of the fluorescence intensity of the bands stained with ethidium bromide in comparison with a DNA size standard of known concentration.

Electrocompetent cells of the $E.\ coli$ K12 strain XL1-Blue (Bullock et al., BioTechniques 5 (1987), 376-379) were prepared according to the methods described by Tung and Chow (Trends Genet. 11 (1995), 128-129) and by Hengen (Trends Biochem. Sci. 21 (1996), 75-76). 1 l of LB medium was adjusted to an optical density at 600 nm, $OD_{600} = 0.08$ by adding a stationary XL1-Blue overnight culture and incubated in a 3 l Erlenmeyer

flask at 200 rpm and 26°C. After reaching $OD_{600} = 0.6$, the culture was cooled on ice for 30 min and then centrifuged at 4000 g and 4°C for 15 min. The cell sediment was washed twice with in each case 500 ml of ice cold 10% w/v glycerol and finally resuspended in 2 ml of ice cold GYT medium (10% w/v glycerol, 0.125% w/v yeast extract, 0.25% w/v tryptone).

Electroporation was carried out by using the Easyjac T Basic system (EquiBio) with the corresponding cuvettes 10 (electrode distance 2 mm). All operational steps were carried out in a cold room at 4°C. 5 to 6 µl of the above-described DNA solution (245 ng/µl) were in each case mixed with 40 µl of the cell suspension, incubated on ice for 1 min and then transferred into the cuvette. 15 After electroporation, the suspension was immediately diluted in 2 ml of fresh ice-cold SOC medium (2% w/v tryptone, 0.5% w/v yeast extract, 10 mM NaCl, 10 mM MgSO₄, 10 mM MgCl₂) and agitated at 37°C and 200 rpm for 60 min. The cells were then sedimented at 3600 g for in 20 each case 2 min, resuspended in 1 ml of LB medium containing 100 µg/ml of ampicillin (LB/Amp) and plated out at 200 µl each on agar plates (140 mm in diameter) with LB/Amp medium. Using a total of 10.7 µg of the 25 ligated DNA in eight electroporation mixtures produced way $3.73 \cdot 10^8$ transformants which this were distributed on 40 agar plates.

incubation 32°C for 14 h, the at colonies After obtained in this way were scraped off the agar plates 30 with the addition of in each case 10 ml of 2xYT/Amp medium, transferred to a sterile Erlenmeyer flask and agitated at 37°C, 200 rpm for 20 min to complete resuspension. 50 ml of 2xYT/Amp medium prewarmed to 37°C were inoculated with 2.88 ml of said suspension so 35 that the cell density was 1.0 OD_{550} . This culture was 37°C, 160 rpm incubated at for 6 h to reach stationary cell density, and phasmid DNA was isolated with the aid of the plasmid Midi kit (Qiagen) according to the manufacturer's instructions. Finally, the DNA was taken up in 100 μl of TE (10 mM Tris/HCl pH 8.0, 1 mM EDTA) and stored at 4°C for further use.

In order to prepare a library of recombinant phagemids (Kay et al., Phage Display of Peptides and Proteins - A Laboratory Manual (1996), Academic Press) which carry the muteins of the bilin-binding protein as a fusion with the truncated coat protein pIII, the phasmid DNA obtained in this way was used for transformation of 10 electrocompetent cells of E. coli Electroporation was carried out as described above with the aid of the Easyjec T Basic system. In a total of 13 mixtures, 40 µl of the cell suspension 15 electrocompetent cells were in each case transformed with in each case 2 µg of the DNA in a volume of 5 µl. After electroporation, the cell suspension obtained from each mixture was diluted immediately in 2 ml of fresh ice-cold SOC medium and agitated at 37°C and 200 rpm for 60 min. 20

The mixtures were combined (volume = 26 ml) and 74 mlof 2xYT medium and 100 µl of ampicillin (stock solution 100 mg/ml, final concentration 100 mg/l) were added. total number of transformants obtained 25 estimated at $1.1 \cdot 10^{10}$ by plating out 100 µl of a $1:10^5$ dilution of the obtained suspension on agar plates containing LB/Amp medium. After incubation at 37°C and 160 rpm for 60 min, the culture was infected with of VCS-M13 helper phage (1.1.1012 pfu/ml, 500 µl 30 Stratagene) and agitated at 37°C, 160 rpm for a further 200 µl of Subsequently, kanamycin solution 35 mg/ml, final concentration 70 mg/l) were added, the incubator temperature was lowered to 26°C 10 min, anhydrotetracyclin (50 µl 35 and, after 50 µg/ml stock solution in dimethylformamide, final concentration $25 \mu g/1)$ was added to induce expression. Finally, the phagemids were produced by incubating the culture at 26°C, 160 rpm for 7 h.

The cells were removed by centrifugation of the culture (15 min, 12000 g, 4°C). The supernatant containing the phagemid particles was sterile-filtered (0.45 μm), mixed with 1/4 volume (25 ml) of 20% w/v PEG 8000, 15% incubated at 4°C overnight. 5 w/v NaCl and centrifugation (20 min, 18000 g, 4°C), the precipitated phagemid particles were dissolved in a total of 4 ml of cold PBS (4 mM KH₂PO₄, 16 mM Na₂HPO₄, 115 mM NaCl, pH 7.4). The solution was incubated on ice for 30 min and distributed into four 1.5 ml reaction vessels at equal 10 After removing undissolved components centrifugation (5 min, 18500 g, 4°C), the supernatant was transferred in each case to a new reaction vessel.

The phagemid particles were again precipitated by mixing with 1/4 volume (in each case 0.25 ml per reaction vessel) of 20% w/v PEG 8000, 15% w/v NaCl and incubating on ice for 60 min. After centrifugation (20 min, 18500 g, 4°C), the supernatant was removed and the precipitated phagemid particles were each dissolved in 0.5 ml of PBS. After incubation on ice for 30 min, centrifugation (5 min, 18500 g, 4°C) was repeated to clarify the solution. The supernatant containing the phagemid particles (between 1·10¹² and 5·10¹² cfu/ml) was then used for affinity concentration.

The recombinant phagemids presenting the muteins of the bilin-binding protein were affinity-concentrated using Immuno Sticks (NUNC). These were coated overnight with 800 μ l of a conjugate (100 μ g/ml) of ribonuclease A (RNaseA) and digoxigenin in PBS.

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1.46 µmol prepared by adding The conjugate was of digoxigenin-3-0-methylcarbonyl-ε-amino-(0.96 mg)acid N-hydroxysuccinimide ester caproic (DIG-NHS, Boehringer Mannheim) in 25 µl of DMSO in µl steps and with constant mixing to 0.73 µmol (10 mg) of RNaseA (Fluka) in 1 ml of 5% w/v sodium hydrogen carbonate. The mixture was incubated with stirring

temperature (RT) for 1 h. Excess reagent was then removed from the RNaseA conjugate by means of a PD-10 gel filtration column (Pharmacia) according to the manufacturer's instructions.

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Unoccupied binding sites on the Immuno Stick surface were saturated by incubation with 1.2 ml of 2% w/v BSA in PBST (PBS with 0.1% v/v Tween 20) at RT for 2 h. After three short washes with in each case 1.2 ml of PBST, the Immuno Stick was incubated in a mixture of 250 μ l of phagemid solution and 500 μ l of blocking buffer (2% w/v BSA in PBST) at RT for 1 h.

Unbound phagemids were removed by stripping off the solution and washing the Immuno Stick eight times with in each case 950 µl of PBST for 2 min. Finally, adsorbed phagemids were competitively eluted during a 15 minute incubation of the Immuno Stick with 950 µl of a 2 mM solution of digoxigenin in PBS (0.742 mg of digoxigenin (Fluka) were to this end dissolved in 19.2 µl of DMF and added to 930.8 µl of PBS).

The phagemids were propagated by heating 950 µl of solution of the elution fraction obtained (between 106 10⁸ colony-forming units, and depending selection cycle) briefly to 37°C, mixing the solution with 4 ml of an exponentially growing culture of E. coli XL1-Blue (OD₅₅₀ = 0.5) and incubated at 200 rpm for 30 min. The phagemid-infected cells were then sedimented (2 min, 4420 g, 4°C), resuspended in 800 μl of fresh 2xxYT medium and plated out on four plates containing LB/Amp medium (140 mm diameter). After incubation at 32°C for 14 h, colonies obtained in this way were scraped off the agar plates with the addition of in each case 10 ml of 2xYT/Amp medium, transferred to a sterile Erlenmeyer flask and agitated at 37°C, 200 rpm for 20 min to complete resuspension.

Production and affinity concentration of phagemid particles were repeated by inoculating 50 ml of 2xYT/Amp medium prewarmed to $37^{\circ}C$ with 0.2 to 1 ml of said suspension so that the cell density was 0.08 OD_{550} . This culture was incubated at $37^{\circ}C$, 160 rpm to a cell density of $OD_{550} = 0.5$, infected with 250 µl of VCS-M13 helper phage (1.1·10¹² pfu/ml, Stratagene), and the procedure was continued as already described above.

from affinity 10 The phagemids obtained the first concentration were used to carry out a series of eight further concentration cycles using Immuno Sticks which had been freshly coated with the digoxigenin-RNaseA The phagemids obtained after conjugate. the last concentration cycle were again used for infecting 15 E. coli XL1-Blue. The mixture of the colonies obtained was scraped off the agar plates using 2xYT/Amp medium described resuspended, as above. This suspension was used to inoculate 50 ml of 2xYT/Amp medium, and the phasmid DNA was isolated using the 20 QIAprep Spin Miniprep kit (QIAGEN) according to the manufacturer's instructions.

In order to be able to produce the muteins of the bilin-binding protein as a fusion protein with the Strep tag II and the albumin-binding domain, the gene cassette between the two BstXI cleavage sites was subcloned from vector pBBP20 into vector pBBP22. A relevant section of the pBBP22 nucleic acid sequence is represented, together with the encoded amino acid sequence, as SEQ ID NO. 9 in the sequence listing. The section starts with the XbaI cleavage site and ends with the HindIII cleavage site. The vector elements outside this region are identical to vector pASK75.

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For this purpose, the DNA isolated from the mixture of the $E.\ coli$ colonies was cut with restriction enzyme BstXI, and the smaller of the two fragments (335 bp) was purified by preparative agarose gel electrophoresis

as described above. In the same manner, pBBP22 vector DNA was cut with BstXI and the larger of the two fragments (3545 bp) was isolated.

- 5 1.5 Weiss units of T4 DNA ligase (Promega) were added to 50 fmol of each of the two DNA fragments in a total volume of 20 μl (30 mM Tris/HCl pH 7.8, 10 mM MgCl₂, 10 mM DTT, 1 mM ATP) and the mixture was incubated for ligation at 16°C overnight. 5 μl of this ligation 10 mixture were used to transform 200 μl of competent cells of E. coli strain TG1-F according to the CaCl₂ method (Sambrook et al., supra), and 2.2 ml of a cell suspension were obtained.
- The transformants were then screened for production of muteins with binding activity for the digoxigenin group by means of a colony screening assay. For this purpose, a cut-to-fit hydrophilic PVDF membrane (Millipore, type GVWP, pore size 0.22 μm) was marked at one position and placed on an LB/Amp agar plate. 150 μl of the cell suspension from the transformation mixture were plated out evenly on said membrane, and approx. 500 colonies were obtained. The plate was incubated in an incubator at 37°C for 7.5 h until the colonies were approx.

In the meantime, a hydrophobic membrane (Millipore, Immobilon P, pore size 0.45 µm) which had likewise been wetted with PBS according to the cut to fit was 30 manufacturer's instructions and subsequently gently agitated in a solution of 10 mg/ml of human serum albumin (HSA, Sigma) in PBS at RT for 4 h. Remaining on the membrane were sites saturated incubation with 3% w/v BSA, 0.5% v/v Tween 20 in PBS at RT for 2 h. The membrane was washed with 20 ml of PBS 35 for two times 10 min and then gently agitated in 10 ml which 200 µg/1 of of LB/Amp medium to anhydrotetracyclin had been added for 10 min. membrane was then marked at one position and placed on a culture plate with LB/Amp agar which additionally contained 200 $\mu g/l$ of anhydrotetracyclin.

The previously obtained hydrophilic membrane on which colonies had grown was then placed onto the hydrophobic membrane such that the two markings coincided. culturing plate with the two membranes was incubated at During this phase, the particular for 15 h. secreted by the colonies as were muteins proteins and immobilized on the lower membrane by means of complex formation between the albumin-binding domain and the HSA.

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the upper membrane containing Subsequently, colonies was transferred to a fresh LB/Amp agar plate 15 stored at 4°C. The hydrophobic membrane removed, washed with 20 ml of PBST for three times 10 min and then incubated in 10 ml of a solution of a conjugate of BSA with digoxigenin in PBST 20 for 1 h.

The conjugate of BSA (Sigma) and digoxigenin was prepared by adding a solution of 3.0 µmol (1.98 mg) of DIG-NHS in 25 µl of DMSO in µl steps and with constant mixing to 300 nmol (19.88 mg) of BSA (Sigma) in 1.9 ml of 5% w/v sodium hydrogen carbonate. The mixture was incubated with stirring at RT for 1 h and excess reagent was removed from the BSA conjugate by means of a PD-10 gel filtration column according to the manufacturer's instructions.

In order to detect bound digoxigenin-BSA conjugate, the membrane was incubated, after washing twice in 20 ml of PBST, with 10 ml of anti-digoxigenin Fab-alkaline phosphatase conjugate (Boehringer Mannheim, diluted 1:1000 in PBST) for 1 h. The membrane was then washed twice with 20 ml PBST and twice with 20 ml of PBST for in each case 5 min and gently agitated in AP buffer (0.1 M Tris/HCl pH 8.8, 0.1 M NaCl, 5 mM MgCl₂) for

10 min. For the chromogenic detection reaction, membrane was incubated in 10 ml of AP buffer to which 5-bromo-4-chloro-3-indolyl phosphate, salt 50 µg/ml toluidinium (BCIP, Roth, in dimethylformamide) and 5 µl of Nitro Blue Tetrazolium (NBT, Sigma, 75 µg/ml in 70% v/v dimethylformamide) had been added, until at the positions of some of the colonies distinct color signals became visible. In this way, digoxigenin-binding activity of the bilin-binding protein muteins which had been produced in the form of fusion proteins with Strep tag and ABD by said colonies was detected.

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Four colonies from the upper membrane, which caused a distinct color signal, were used for preparing cultures 15 in LB/Amp medium of 4 ml in volume. Their plasmid DNA isolated with the aid of the JETquick Plasmid kit (Genomed) according to Miniprep Spin manufacturer's instructions, and the gene section 20 coding for the mutein was subjected to sequence analysis. Sequence analysis was carried out with the aid of the T7 sequencing kit (Pharmacia) according to instructions manufacturer's by oligodeoxynucleotides SEQ ID NO. 10 and SEQ ID NO. 11. 25 It was found in the process that all four plasmids studied carried the same nucleotide sequence. corresponding gene product was denoted by DigA (SEQ ID 12). The DigA nucleotide sequence was translated into the amino acid sequence and is represented in the 30 sequence listing.

Example 2

Partial random mutagenesis of the DigA mutein and selection of muteins with improved binding affinity for digoxigenin

In order to improve the affinity between the DigA mutein and digoxigenin, which was determined as

 295 ± 36 nM according to Example 3, the 6 amino acid positions 28, 31 and 34-37 in DigA were selected for a more substantial partial random mutagenesis.

Said positions were mutated by carrying out the PCR using a degenerated oligodeoxynucleotide primer. amplification reaction was carried out in a total 100 µl, with 2 ng of the vector pBBP22 volume of plasmid DNA coding for DigA (SEQ ID NO. 12) being used 10 as matrix. The reaction mixture contained 50 pmol of the two primers SEQ ID NO. 13 and SEQ ID NO. 7 and also the other components according to the method described in Example 1. The PCR was carried out in 20 temperature cycles of 1 min at 94°C, 1 min at 65°C, and 1.5 min at 15 72°C, followed by a final incubation at 60°C for 5 min. The DNA fragment obtained was isolated by preparative agarose gel electrophoresis and then cut with BstXI according the manufacturer's instructions. to resulting DNA fragment of 335 bp in length was again 20 purified by preparative agarose gel electrophoresis.

PBBP24 vector DNA was cut with BstXI accordingly and the 4028 bp fragment obtained was isolated. A relevant section of the pBBP24 nucleic acid sequence 25 represented, together with the encoded amino acid sequence, as SEQ ID NO. 14 in the sequence listing. The section starts with the XbaI cleavage site and ends with the *Hind*III cleavage site. The vector elements outside this region are identical to vector pASK75. PBBP24 is virtually identical with pBBP20, and the BBP 30 gene has been inactivated by means of appropriately introduced stop codons.

1.3 μg of the cleaved DNA fragment from the PCR and 16.0 μg of the pBBP24 fragment were incubated for ligation in the presence of 120 Weiss units of T4 DNA ligase (New England Biolabs) in a total volume of 600 μl (50 mM Tris/HCl pH 7.8, 10 mM MgCl₂, 10 mM DTT, 1 mM ATP, 50 $\mu g/m l$ BSA) at 16°C for 18 h. The DNA was

then precipitated by adding 10 μg of yeast tRNA (Boehringer Mannheim), 25 μl of 5 M ammonium acetate and 100 μl of ethanol to in each case 24 μl of the ligation mixture. Incubation at -20°C for two weeks was followed by centrifugation (20 min, 16000 g, 4°C). The precipitate was washed in each case with 150 μl of ethanol (70% v/v, -20°C) and dried under reduced pressure. Finally, the DNA was taken up in 80 μl of TE/10.

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- E . coli XL1-Blue cells were transformed with ligated DNA by electroporation according to procedure described in Example 1, with in each case 40 µl of cell suspension of electrocompetent cells 15 being mixed with 5 µl of the DNA solution in mixtures. After electroporation, the cells were immediately diluted in 2 ml of fresh ice-cold SOC medium and agitated at 37°C and 200 rpm for 60 min.
- 168 ml of 2xYT medium and 200 µl of ampicillin (stock 20 solution 100 mg/ml, final concentration 100 mg/l) were added to the combined suspensions. The total number of transformants obtained was estimated at 1.48•109 by plating out 100 µl of a 1:104 dilution of the obtained 25 cell suspension on agar plates. After incubation at 37°C and 160 rpm for 60 min, the transformants were infected with 4 ml of VCS-M13 helper phage (6.3.1011 pfu/ml, Stratagene) and agitated at 37°C and 160 rpm for a further 30 min. Subsequently, 400 µl of kanamycin 30 (stock solution 35 mg/ml, final concentration 70 mg/l) were added, the incubator temperature was lowered to 26°C and, after 10 min, anhydrotetracyclin (100 μl of a 50 µg/ml stock solution in dimethylformamide, concentration $25 \mu g/1)$ was added to induce 35 expression. Finally, the phagemids were produced by incubating the culture at 26°C and 160 rpm for 7 h. The cells were removed and the phagemids purified by precipitation as described in Example 1.

Streptavidin-coated paramagnetic particles (Dynabeads M-280 Streptavidin, Dynal) were used together with a double conjugate of BSA with digoxigenin and biotin for affinity concentration from the library of phagemids which presented the partially mutated DigA mutein.

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A double conjugate of BSA with digoxigenin and biotin was prepared by adding 1.5 μ mol (0.99 mg) of DIG-NHS in and 1.5 µmol (0.68 mq)12.5 ul of DMSO of D-biotinoyl-&-aminocaproic acid N-hydroxysuccinimide 10 ester (Boehringer Mannheim) in 12.5 μl of DMSO in μl steps and with constant mixing to 300 nmol (19.88 mg) of BSA in 1.9 ml of 5% w/v sodium hydrogen carbonate. The mixture was incubated with stirring at RT for 1 h. Excess reagent was removed from the double conjugate 15 via a PD-10 gel filtration column according to the manufacturer's instructions.

Digoxigenin-binding phagemids were concentrated 20 mixing 40 µl of a 0.5 µM solution of the double conjugate (33.5 μ g/ml) in PBS with 260 μ l of a solution of the freshly prepared phagemids (between 5.1011 and 5.10¹² cfu/ml) and incubated at RT for 1 h so that the complex between the digoxigenin group and the muteins presented by the phagemids was able to form. This was 25 followed by adding 100 µl of a solution of 8% w/v BSA, 0.4% v/v Tween 20 in PBS.

Parallel thereto, 100 µl of the commercially available suspension of paramagnetic particles were washed with three times 100 µl of PBS. Here, the particles were suspended for 1 min by rotating the Eppendorf vessel and then collected at the wall of the Eppendorf vessel with the aid of a magnet, and the supernatant was stripped off. Unspecific binding sites 35 were saturated by incubating the paramagnetic particles with 100 μ l of 2% w/v BSA in PBST at RT for 1 h. After removing the supernatant, the mixture of conjugate and phagemids was added to the paramagnetic particles, and the particles were resuspended and incubated at RT for 10 min. Finally, free biotin-binding sites of Streptavidin were saturated by adding 10 μ l of a 4 μ M D-dethiobiotin (Sigma) solution in PBS to the mixture and incubating said mixture at RT for 5 min. This procedure also prevented the Strep tag II as part of the fusion protein of the muteins and the phage coat protein pIII fragment from being able to form a complex with Streptavidin.

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Unbound phagemids were removed by washing paramagnetic particles with eight times 1 ml of fresh PBST with the addition of 1 mM D-dethiobiotin, particles were collected with the aid of the magnet and the supernatant was stripped off. The bound phagemids were eluted by incubating the resuspended particles in 950 µl of 0.1 M glycine/HCl pH 2.2 for 15 minutes. After collecting the particles on the magnet, again stripped off and this supernatant was was immediately followed by neutralizing the pH of said solution by addition of 140 μ l of 0.5 M Tris.

The phagemids were propagated by mixing the elution fraction obtained, according to the procedure Example 1, with 4 ml of an exponentially growing culture of $E.\ coli\ XL1-Blue\ (OD_{550}=0.5)$ and incubating 37°C, 200 rpm for 30 min. The phagemid-infected cells were then sedimented (2 min, 4420 g, resuspended in 800 µl of fresh 2xYT medium and plated on four agar plates containing LB/Amp medium out (140 mm in diameter). After incubation at 32°C for 14 h, the colonies obtained in this way were scraped off the agar plates with the addition of in each case 10 ml of 2xYT/Amp medium, transferred to a sterile Erlenmeyer flask and agitated at 37°C, 200 rpm for 20 min to complete resuspension.

Production and affinity concentration of phagemid particles were repeated by inoculating 50 ml of

2xYT/Amp medium prewarmed to 37°C with 0.2 to 1 ml of said suspension so that the cell density was 0.08 OD₅₅₀. This culture was incubated at 37°C, 160 rpm to a cell density of OD_{550} = 0.5 and infected with 300 μl of VCS-M13 helper phage $(6.3 \cdot 10^{11} \text{ pfu/ml, Stratagene})$. affinity selection was then repeated particles and the digoxigenin/biotin paramagnetic double conjugate under the abovementioned conditions. A total of 4 selection cycles were carried out in this way.

The phagemids obtained after the last concentration cycle were again used for infecting *E. coli* XL1-Blue. The mixture of the obtained colonies which had been scraped off the agar plates using 2xYT/Amp medium and had been resuspended, as described above, was used to inoculate 50 ml of 2xYT/Amp medium, and phasmid DNA was isolated using the QIAprep spin miniprep kit (QIAGEN) according to the manufacturer's instructions.

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Subsequently, the gene cassette between the two BstXI cleavage sites was subcloned, as in Example 1, from vector pBBP24 into vector pBBP22, and competent cells of E. coli strain TG1-F were transformed according to the CaCl2 method. Finally, the transformants were, again according to Example 1, screened for production of muteins with binding activity for the digoxigenin group by means of the colony screening assay.

Seven of the colonies showing a strong signal intensity 30 in the colony screening assay were cultured. plasmid DNA was isolated with the aid of the plasmid (Genomed) according spin kit manufacturer's instructions, and the gene subjected to coding for the mutein was sequence analysis as in Example 1. It was found in the process 35 that all plasmids studied had different sequences. After translating the nucleotide sequences into amino acid sequences, six of the seven variants studied had an amber stop codon at amino acid position 28. However,

this stop codon was at least partially suppressed when choosing suitable amber-suppressor strains such as, for example, $E.\ coli$ XL1-Blue or TG1-F and instead translated as glutamine. Thus a full-length functional protein was produced both during affinity concentration and in the colony screening assay.

As the only mutein without an amber stop codon among the muteins found, the mutein with SEQ ID NO. 15 was particularly well suited to bacterial production. Consequently, this mutein, also denoted by DigA16, was characterized in more detail with regard to its ability to bind to the digoxigenin group.

15 Example 3

Production of the DigA and DigA16 muteins and determination of their affinity for digoxigenin and derivatives thereof by fluorescence titration

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The bilin-binding protein muteins obtained from the previous examples were preparatively produced by subcloning the coding gene section between the two BstXI cleavage sites from the type pBBP22 vector into the expression plasmid pBBP21. The plasmid thus obtained coded for a fusion protein of the OmpA signal sequence, followed by the mutein and the Strep tag II affinity tag.

30 A relevant section of the pBBP21 nucleic acid sequence is represented, together with the encoded amino acid sequence, as SEQ ID NO. 16 in the sequence listing. The section starts with the XbaI cleavage site and ends with a hexanucleotide which was obtained by ligating a blunt strand end with a filled-up HindIII strand end, with the loss of the original HindIII cleavage site. The vector elements outside this region are identical to vector pASK75.

For subcloning, the plasmid DNA coding for the relevant mutein was cut with restriction enzyme BstXI, and the smaller of the two fragments (335 bp) was purified by preparative agarose gel electrophoresis as described in Example 1. In the same manner, pBBP21 vector DNA was cut with BstXI, and the larger of the two fragments (4132 bp) was isolated.

1.5 Weiss units of T4 DNA ligase (Promega) were added to 50 fmol of each of the two DNA fragments in a total volume of 20 μl (30 mM Tris/HCl pH 7.8, 10 mM MgCl₂, 10 mM DTT, 1 mM ATP) and the mixture was incubated for ligation at 16°C for 16 h. 5 μl of the ligation mixture were then used to transform E. coli JM83 (Yanisch-Perron et al., Gene 33 (1985), 103-119) according to the CaCl₂ method, and 2.2 ml of a cell suspension were obtained. 100 μl of this suspension were plated out on an agar plate containing LB/Amp medium and incubated at 37°C for 14 h.

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The protein was produced by selecting one of the obtained single colonies, using it to inoculate a 50 ml preculture (LB/Amp medium) and incubating said preculture at 30°C and 200 rpm overnight. 40 ml of the preculture were then transferred by inoculating 2 l of LB/Amp medium in a 5 l Erlenmeyer flask, followed by incubating the culture at 22°C and 200 rpm. At a cell density of $OD_{550} = 0.5$, gene expression was induced by adding 200 µg/l anhydrotetracyclin (200 µl of a 2 mg/ml stock solution in DMF), followed by agitating at 22°C, 200 rpm for a further 3 h.

The cells were removed by centrifugation (15 min, 4 420 g, 4°C) and, after removing the supernatant, resuspended in 20 ml of periplasm lysis buffer (100 mM Tris/HCl pH 8.0, 500 mM sucrose, 1 mM EDTA) with cooling on ice. After incubation on ice for 30 min, the spheroplasts were removed in two successive centrifugation steps (15 min, 4 420 g, 4°C and 15 min,

30 000 g, 4° C). The periplasmic protein extract obtained in this way was dialyzed against SA buffer (100 mM Tris/HCl pH 8.0, 150 mM NaCl, 1 mM EDTA), sterile-filtered and used for chromatographic purification.

Purification was carried out by means of the Strep tag II affinity tag (Schmidt and Skerra, Protein Eng. 6 (1993), 109-122) fused to the C terminus of the muteins. In the present case, Streptavidinmutein "1" was used (Voss and Skerra, Protein Eng. 10 (1997), 975-982), which was coupled to activated Sepharose (via 5 mg/ml immobilized Streptavidin, based on the bett volume of the matrix).

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A chromatography column packed with 2 ml of said material was equilibrated at 4°C and a flow rate of 20 ml/h with 10 ml of SA buffer. The chromatography was monitored by measuring absorption of the eluate at 280 nm in a flow-through photometer. Application of the periplasmic protein extract was followed by washing with SA buffer until the base line was reached. Bound mutein was then eluted with 10 ml of a solution of 2.5 mM D-dethiobiotin (Sigma) in SA buffer. The fractions containing the purified mutein were checked by means of SDS polyacrylamide gel electrophoresis (Fling and Gregerson, Anal. Biochem. 155 (1986), 83-88) and combined. The protein yields were between 200 µg and 800 µg per 2 l culture.

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The ligand binding properties of muteins DigA, DigA16 and also of the recombinant bilin-binding protein (SEQ ID NO: 16) were determined by means of the method of fluorescence titration.

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In this case, the decrease in intrinsic tyrosine and/or tryptophan fluorescence of the protein forming a complex with the ligand was measured. The measurements were carried out in a fluorimeter, type LS 50 B (Perkin

Elmer) at an excitation wavelength of 295 nm (slit width 4 nm) and an emission wavelength of 345 nm (slit width 6 nm). The ligands used were digoxigenin (Fluka), digoxin (Fluka), digitoxigenin (Fluka), digitoxin (Fluka), testosterone (Sigma), ouabain (Fluka), and 4-aminofluorescein (Fluka). The ligands showed no significant intrinsic fluorescence or absorption at the stated wavelength.

The buffer system used was PBS with the addition of 10 1 mM EDTA. The solution of the relevant purified mutein dialyzed four times against this buffer adjusted to a concentration of 1 µm by dilution. All solutions used were sterile-filtered (Filtropur $0.45 \mu m$, Sarstedt). The concentration was determined by 15 280 nm using calculated absorption at of extinction coefficients of 53 580 M⁻¹ cm⁻¹ for DigA and DigA16 (Wisconsin Software Package, Genetics Computer Group). For Bbp, the calculated extinction coefficient $54 150 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1}$, corrected in the presence of 20 guanidinium chloride according to Gill and von Hippel (Anal. Biochem. 182 (1989), 319-326) was used.

For the measurement, 2 ml of the mutein solution were introduced into a quartz cuvette equipped with a 25 magnetic stirrer bar and thermally equilibrated at 25°C in the sample holder of the photometer. Then a total of 40 µl of a 100 µM to 500 µM solution of the ligand in the same buffer were pipetted in steps of from $1 \mu l$ to 30 4 µl. The dilution of the introduced protein solution by altogether no more than 2%, which took place in the process, was not taken into account in the subsequent evaluation of the data. After each titration step, the equilibrium was allowed to form by incubating with stirring for 1 min, and the fluorescence signal was 35 measured as average over 10 s. After subtracting the fluorescence value of the buffer, the signals were normalized to an initial value of 100%.

The thus obtained data of a titration series were fitted by nonlinear regression with the aid of the computer program Kaleidagraph (Abelbeck Software) according to the following formula

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$$F = ([P]_t - [L]_t - K_d) \frac{f_P}{2} + ([P]_t + [L]_t + K_d) \frac{f_{PL}}{2} + (f_P - f_{PL}) \sqrt{\frac{([P]_t + [L]_t + K_d)^2}{4} - [P]_t [L]_t}$$

Here, F means the normalized fluorescence intensity and $[L]_t$ the total ligand concentration in the particular titration step. $[P]_t$ as mutein concentration, f_{PL} as fluorescence coefficient of the mutein-ligand complex and K_d as the thermodynamic dissociation constant of said complex were fitted as free parameters to the normalized data.

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Fig. 1 represents graphically the results of the fluorescence titrations of the DigA16 mutein with the ligands digoxigenin, digitoxigenin and ouabain. It turns out that digitoxigenin is bound even tighter than digoxigenin, while no binding is observed for ouabain.

The values resulting from the fluorescence titrations for the dissociation constants of the complexes of the bilin-binding protein muteins and the various ligands are summarized in the following table:

	Bbp variant	Ligand	K_{d} [nM]
	Bbp:	digoxigenin	_*
30	DigA:	digoxigenin	295 ± 37
		digoxin	200 ± 34
	DigA16:	digoxigenin	30.2 ± 3.6
	_	digoxin	31.1 ± 3.2
35		digitoxigenin	2.8 ± 2.7
		digitoxin	2.7 ± 2.0
		ouabain	_*

testosterone -*
4-aminofluroescein -*

no detectable binding activity

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Example 4

Preparation of fusion proteins of the DigA16 mutein and bacterial alkaline phosphatase and use for detecting digoxigenin groups in an ELISA and in a Western blot

In order to produce two different fusion proteins of the DigA16 mutein and bacterial alkaline phosphatase (PhoA) with different arrangement of the partners within the polypeptide chain, the two expression plasmids pBBP27 and pBBP29 were constructed by using the molecular-biological methods familiar to the skilled worker.

pBB27 codes for a fusion protein of PhoA including the signal sequence thereof, a short peptide linker having the amino acid sequence Pro-Pro-Ser-ala, the sequence corresponding to the mature DigA16 mutein and the Strep tag II. A relevant section of the pBBP27 nucleic acid sequence is represented, together with the encoded amino acid sequence, as SEQ ID NO: 17 in the sequence listing. The section begins with the XbaI cleavage site and ends with the HindIII cleavage site. The vector elements outside this region are identical to vector pBBP21.

pBB29 codes for a fusion protein of DigA16 with preceding OmpA signal sequence, followed by the peptide sequence for Strep tag II, a sequence of 5 glycine residues and the mature PhoA sequence without the N-terminal amino acid arginine. A relevant section of the pBBP29 nucleic acid sequence is represented, together with the encoded amino acid sequence, as SEQ ID NO: 18 in the sequence listing. The section begins with the

XbaI cleavage site and ends with the HindIII cleavage site. The vector elements outside this region are identical to vector pBBP21.

- 5 Both plasmids additionally code for the bacterial protein disulfide isomerase DsbC on a separate cistron located in 3' direction. The plasmids are shown diagrammatically in Fig. 2.
- The fusion proteins encoded by plasmids pBBP27 and 10 pBBP29 were produced analogously to the method for preparing the simple muteins, described in example 3. In order to avoid complexing the metal ions from the active center of PhoA, lysis of the bacterial periplasm 15 using EDTA-free carried out lysis Polymyxin B sulfate (2 mg/ml, Sigma) was added to the buffer as an agent destabilizing the outer cell membrane. All other buffers used for purification were likewise EDTA-free.

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The fusion proteins purified by affinity chromatography by means of the Strep tag II were dialyzed against PBS buffer overnight. The fusion protein yields between 100 and 200 µg per 2 l of culture medium. The purity of the fusion proteins obtained was checked by SDS polyacrylamide gel electrophoresis, according to example 3, and determined at 90-95%. Subsequently, the fusion proteins were used for directly detecting the digoxigenin group conjugates of with proteins both in a sandwich ELISA and in a Western blot.

While the conjugates used of digoxigenin with RNaseA and BSA were prepared according to example 1, a conjugate of digoxigenin with ovalbumin (Sigma) was prepared by adding 1.5 µmol (0.99 mg) DIG-NHS in 25 µl of DMSO in µl steps and with constant mixing to 300 nmol (13.5 mg) of ovalbumin in 1.9 ml of 5% sodium hydrogen carbonate. The mixture was incubated with

stirring at RT for 1 h. Excess reagent was removed from the ovalbumin conjugate via a PD-10 gel filtration column according to the manufacturer's instructions.

For detecting digoxigenin groups in a sandwich ELISA, the wells in in each case two vertical columns of a microtiter plate (ELISA strips, 2×8 well with high binding capacity, F type, Greiner) were filled in each case with 100 μ l of a 100 μ g/ml solution of the BSAdigoxigenin conjugate or the ovalbumin-digoxigenin 10 conjugate in PBS and incubated at RT overnight. As a control, the wells of a fifth vertical row of the microtiter plate were filled with 100 µl of a 100 µg/ml solution of nonconjugated BSA (Sigma) PBS in 15 likewise incubated at RT overnight. After removing the solution, unoccupied binding sites were saturated with 200 μ l of a solution of 2% w/v BSA in PBST for 2 h. After washing three times with PBST, 50 μ l of a 1 μ M solution of the purified fusion protein were in each 20 case introduced into the first well of a row, and the Tween concentration was adjusted to 0.1% v/v by adding 1 μl of a solution of 5% v/v Tween. The subsequent wells in each row were initially charged with 50 µl of PBST. Then, 50 µl of the purified fusion protein were 25 pipetted in each case into the second well, mixed and, therefrom, 1:2 dilutions starting were prepared stepwise in the other wells of the vertical row. After incubation at RT for 1 h, the wells were washed twice with PBST and twice with PBS. The fusion proteins bound 30 to the digoxigenin groups were finally detected by means of alkaline phosphatase-catalyzed hydrolysis of p-nitrophenyl phosphate. For this purpose, 100 μ l of a solution of 0.5 mg/ml p-nitrophenyl phosphate (Amresco) in AP buffer (100 mM NaCl, 5 mM MgCl₂, 100 mM Tris/HCl pH 8.8) were introduced into the wells and product 35 formation was monitored by measuring absorption SpectraMax 250 photometer (Molecular 405 nm in a Devices).

Fig. 3 shows the result of this measurement. According to this, the digoxigenin group is recognized both as conjugate with BSA and as conjugate with ovalbumin, leading to the conclusion that binding by the DigA16 mutein is context-independent. Furthermore, both fusion proteins are active both with regard to the binding function for the digoxigenin group and enzymatically and produce, despite their different structure, almost identical signals.

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The fusion proteins encoded by vectors pBBP27 were used in а Western blot by fractionating 5 µl of a protein mixture in PBS, whose of digoxigenin-BSA concentration conjugate, digoxigenin-ovalbumin conjugate and digoxigenin-RNaseA 15 conjugate was simultaneously in each case 100 µg/ml, and also $5\ \mu l$ of a protein mixture in PBS, whose concentration of underivatized BSA, ovalbumin likewise simultaneously RNaseA was in each case 20 100 μg/ml, by SDS polyacrylamide gel electrophoresis. mixture was then transferred The protein nitrocellulose by electrotransfer (Blake et al., Anal. Biochem. 136 (1984), 175-179). The membrane was then washed in 10 ml of PBST for three times 5 min and incubated with 10 ml of a 0.5 μM solution of in each 25 case one of the two fusion proteins for 1 h. membrane was then washed in 10 ml PBST for two times 5 min and in 10 ml of PBS for two times 5 min finally gently agitated in 10 ml of AP buffer 30 10 min. For the chromogenic detection reaction, membrane was incubated in 10 ml of AP buffer to which 30 µl BCIP (50 µg/ml in dimethylformamide) and 5 µl NBT (75 µg/ml in 70% v/v dimethylformamide) had been added, and in this way bound fusion protein was detected.

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Fig. 4 shows the result of this detection method. It turns out again that binding of the digoxigenin group by the two fusion proteins is independent of the carrier protein and that both fusion proteins achieve

comparable signal intensities. The same carrier proteins cause no signal whatsoever if they are not conjugated with the digoxigenin group.

SEQUENCE LISTING

GENERAL INFORMATION:

APPLICANT:

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CITY: Freising COUNTRY: Germany

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TELEPHONE: 08161-714351 TELEFAX: 08161-714352

TITLE OF THE INVENTION: Muteins of the bilin-binding protein

NUMBER OF SEQUENCES: 18

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk COMPUTER: IBM PC compatible OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: Microsoft Word, format: Text

CURRENT APPLICATION DATA:

APPLICATION NUMBER: not yet known

FILING DATE: not yet known

INFORMATION FOR SEQ ID NO:1:

SEQUENCE CHARACTERISTICS:

LENGTH: 1219 base pairs TYPE: nucleic acid STRANDEDNESS: double

TOPOLOGY: linear

MOLECULE TYPE: fragment of phasmid pBBP20

FEATURE:

NAME/KEY: signal peptide

LOCATION: (22..84)

FEATURE:

NAME/KEY: mature peptide LOCATION: (85..1209)

OTHER INFORMATION:

/Product = "fusion protein composed of bilin-binding protein

Strep-tag II and fragment of phage envelope

protein pIII"

/Codon = (sequence: "TAG", amino acid:Gln)

FEATURE:

NAME/KEY: coding sequence

LOCATION: (85..606) OTHER INFORMATION:

/Product = "mature bilin-binding protein"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (607..636)
OTHER INFORMATION:

/Product = "Strep-tag II affinity tag"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (637..639) OTHER INFORMATION:

/Other = "amber stop codon"

FEATURE:

NAME/KEY: coding sequence LOCATION: (640..1209) OTHER INFORMATION:

/Product = "amino acids 217-406 of envelope protein pIII"

SEQUENCE DESCRIPTION: SEQ ID NO:1:

TCTAGTTAAC GAGGGCAAAA A ATG AAA AAG ACA GCT ATC GCG ATT

Met Lys Lys Thr Ala Ile Ala Ile

-21 -20 -15

GCA GTG GCA CTG GCT GGT TTC GCT ACC GTA GCG CAG GCC GAC GTG 90 Ala Val Ala Leu Ala Gly Phe Ala Thr Val Ala Gln Ala Asp Val -10 -5 -1 1

TAC CAC GAC GGT GCC TGT CCC GAA GTC AAG CCA GTC GAC AAC TTC 135
Tyr His Asp Gly Ala Cys Pro Glu Val Lys Pro Val Asp Asn Phe
5 10 15

GAC TGG TCC CAG TAC CAT GGT AAA TGG TGG GAA GTC GCC AAA TAC 180 Asp Trp Ser Gln Tyr His Gly Lys Trp Trp Glu Val Ala Lys Tyr 20 25 30

CCC AAC TCA GTT GAG AAG TAC GGA AAG TGC GGA TGG GCT GAG TAC 225 Pro Asn Ser Val Glu Lys Tyr Gly Lys Cys Gly Trp Ala Glu Tyr 35 40 45

ACT CCT GAA GGC AAG AGT GTC AAA GTT TCG AAC TAC CAC GTA ATC 270 Thr Pro Glu Gly Lys Ser Val Lys Val Ser Asn Tyr His Val Ile 50 55 60

CAC GGC AAG GAA TAC TTT ATT GAA GGA ACT GCC TAC CCA GTT GGT 315 His Gly Lys Glu Tyr Phe Ile Glu Gly Thr Ala Tyr Pro Val Gly 65 70 75

GAC TCC AAG ATT GGA AAG ATC TAC CAC AGC CTG ACT TAC GGA GGT 360 Asp Ser Lys Ile Gly Lys Ile Tyr His Ser Leu Thr Tyr Gly Gly 80 85 90

GTC ACC AAG GAG AAC GTA TTC AAC GTA CTC TCC ACT GAC AAC AAG 405 Val Thr Lys Glu Asn Val Phe Asn Val Leu Ser Thr Asp Asn Lys 95 100 105

AAC Asn	TAC Tyr	Ile 110	Ile	GGA Gly	TAC Tyr	TAC Tyr	TGC Cys 115	AAA Lys	TAC Tyr	GAC Asp	GAG Glu	GAC Asp 120	AAG Lys	AAG Lys	450
GGA Gly	CAC His	CAA Gln 125	GAC Asp	TTC Phe	GTC Val	TGG Trp	GTG Val 130	CTC	TCC Ser	AGA Arg	AGC Ser	ATG Met 135	GTC Val	CTT Leu	495
ACT Thr	GGT Gly	GAA Glu 140	GCC Ala	AAG Lys	ACC Thr	GCT Ala	GTC Val 145	GAG Glu	AAC Asn	TAC Tyr	CTT Leu	ATC Ile 150	GGC Gly	TCC Ser	540
CCA Pro	GTA Val	GTC Val 155	GAC Asp	TCC Ser	CAG Gln	AAA Lys	CTG Leu 160	GTA Val	TAC Tyr	AGT Ser	GAC Asp	TTC Phe 165	TCT Ser	GAA Glu	585
GCC Ala	GCC Ala	TGC Cys 170	AAG Lys	GTC Val	AAC Asn	AAT Asn	AGC Ser 175	Asn	TGG Trp	TCT Ser	CAC His	CCG Pro 180	CAG Gln	TTC Phe	630
GAA Glu	AAA Lys	TAG Gln 185	GCT Ala	GGC Gly	GGC Gly	GGC Gly	TCT Ser 190	GGT Gly	GGT Gly	GGT Gly	TCT Ser	GGC Gly 195	GGC Gly	GGC Gly	675
TCT Ser	GAG Glu	GGT Gly 200	GGT Gly	GGC Gly	TCT Ser	GAG Glu	GGT Gly 205	GGC Gly	GGT Gly	TCT Ser	GAG Glu	GGT Gly 210	GGC Gly	GGC Gly	720
Ser	Glu	Gly 215	Gly	GGT Gly	Ser	Gly	Gly 220	Gly	Ser	Gly	Ser	Gly 225	Asp	Phe	
GAT Asp	TAT Tyr	GAA Glu 230	AAG Lys	ATG Met	GCA Ala	AAC Asn	GCT Ala 235	AAT Asn	AAG Lys	GGG Gly	GCT Ala	ATG Met 240	ACC Thr	GAA Glu	810
AAT Asn	GCC Ala	GAT Asp 245	GAA Glu	AAC Asn	GCG Ala	CTA Leu	CAG Gln 250	TCT Ser	GAC Asp	GCT Ala	AAA Lys	GGC Gly 255	AAA Lys	CTT Leu	855
GAT Asp	TCT Ser	GTC Val 260	GCT Ala	ACT Thr	GAT Asp	TAC Tyr	GGT Gly 265	GCT Ala	GCT Ala	ATC Ile	GAT Asp	GGT Gly 270	TTC Phe	ATT Ile	900
GGT Gly	GAC Asp	GTT Val 275	TCC Ser	GGC Gly	CTT Leu	GCT Ala	AAT Asn 280	GGT Gly	AAT Asn	GGT Gly	GCT Ala	ACT Thr 285	GGT Gly	GAT Asp	945
TTT Phe	GCT Ala	GGC Gly 290	TCT Ser	AAT Asn	TCC Ser	CAA Gln	ATG Met 295	GCT Ala	CAA Gln	GTC Val	GGT Gly	GAC Asp 300	GGT Gly	GAT Asp	990
				ATG Met											1035

		DE 199	26 068	C1		
Pro Gln S	CCG GTT GAA T Ser Val Glu C 320	GT CGC CCT ys Arg Pro 325	Phe Val	TTT GGC Phe Gly	GCT GGT Ala Gly 330	AAA 1080 Lys
Pro Tyr G	GAA TTT TCT A Glu Phe Ser I 335	TT GAT TGT le Asp Cys 340	Asp Lys	ATA AAC Ile Asn	TTA TTC Leu Phe 345	CGT 1125 Arg
Gly Val P	TTT GCG TTT C Phe Ala Phe L 350	TT TTA TAT eu Leu Tyr 355	Val Ala	ACC TTT Thr Phe	ATG TAT Met Tyr 360	GTA 1170 Val
Phe Ser T	ACG TTT GCT A Thr Phe Ala A 165	AC ATA CTG sn Ile Leu 370	Arg Asn	AAG GAG Lys Glu	TCT Ser 375	1209
TAATAAGCT	T					1219
INFORMATIC	ON FOR SEQ ID	NO:2:				
LENG TYPE STRA TOPO MOLE SEQUENC	CE CHARACTERI STH: 64 bases E: nucleic ac ANDEDNESS: si DLOGY: linear ECULE TYPE: s	id ngle ynthetic	NO:2			
CCATGGTAA ACGGAAAGT	A TGGTGGGAAG G CGGA	TCGCCAAAT	'A CCCCNN	KNMS NNS	NNKAAGT	50 64
INFORMATIC	ON FOR SEQ ID	NO:3:				
LENG TYPE STRA TOPO MOLE	CE CHARACTERI GTH: 71 bases E: nucleic ac ANDEDNESS: si DLOGY: linear ECULE TYPE: s	id ngle ynthetic	-	xynucleo	tide	
GGGTAGGCG	G TACCTTCSNN T TTGACACTCT	AAAGTATTO		TGGA TTA	CMNNGTA	50 71

INFORMATION FOR SEQ ID NO:4:

SEQUENCE CHARACTERISTICS:

LENGTH: 74 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:4:

CCAAGATTGG AAAGATCTAC CACAGCNNSA CTNNKGGAGG TNNSACCVVS 50
GAGNNKGTAT TCAACGTACT CTCC 74

INFORMATION FOR SEQ ID NO:5:

SEQUENCE CHARACTERISTICS:

LENGTH: 78 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:5:

TCTGGAGAGC ACCCAGACMN NGTCSNNGTG TCCCTTCTTG TCCTCGTCGT 50
ASNNGCAMNN GTATCCGATG ATGTAGTT 78

INFORMATION FOR SEQ ID NO:6:

SEQUENCE CHARACTERISTICS:

LENGTH: 36 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:6:

CTTCGACTGG TCCCAGTACC ATGGTAAATG GTGGGA 36

INFORMATION FOR SEQ ID NO:7:

SEQUENCE CHARACTERISTICS:

LENGTH: 37 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:7:

CACCAGTAAG GACCATGCTT CTGGAGAGCA CCCAGAC 37

INFORMATION FOR SEQ ID NO:8:

SEQUENCE CHARACTERISTICS:

LENGTH: 46 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:8:

AGATCTTTCC AATCTTGGAG TCACCAACTG GGTAGGCGGT ACCTTC 46

INFORMATION FOR SEQ ID NO:9:

SEQUENCE CHARACTERISTICS:

LENGTH: 793 base pairs TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: fragment of plasmid pBBP22

FEATURE:

NAME/KEY: signal peptide

LOCATION: (22..84)

FEATURE:

NAME/KEY: mature peptide

LOCATION: (85..783) OTHER INFORMATION:

/Product = "fusion protein composed of bilin-binding

protein, Strep-tag II and albumin-binding

domain"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (85..606) OTHER INFORMATION:

/Product = "mature bilin-binding protein"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (607..636) OTHER INFORMATION:

/Product = "Strep-tag II affinity tag"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (637..783) OTHER INFORMATION:

/Product = "albumin-binding domain of protein G"

SEQUENCE DESCRIPTION: SEQ ID NO:9:

TCTAGATAAC GAGGGCAAAA A ATG AAA AAG ACA GCT ATC GCG ATT

Met Lys Lys Thr Ala Ile Ala Ile

-21 -20 -15

GCA GTG GCA CTG GCT GGT TTC GCT ACC GTA GCG CAG GCC GAC GTG 90
Ala Val Ala Leu Ala Gly Phe Ala Thr Val Ala Gln Ala Asp Val
-10 -5 -1 1

TAC CAC GAC GGT GCC TGT CCC GAA GTC AAG CCA GTC GAC AAC TTC 135
Tyr His Asp Gly Ala Cys Pro Glu Val Lys Pro Val Asp Asn Phe
5 10 15

GAC Asp	TGG Trp	TCC Ser 20	CAG Gln	TAC Tyr	CAT	GGT Gly	AAA Lys 25	TGG Trp	TGG Trp	GAA Glu	GTC Val	GCC Ala 30	AAA Lys	TAC Tyr	180
CCC Pro	AAC Asn	TCA Ser 35	GTT Val	GAG Glu	AAG Lys	TAC Tyr	GGA Gly 40	AAG Lys	TGC Cys	GGA Gly	TGG Trp	GCT Ala 45	GAG Glu	TAC Tyr	225
ACT Thr	CCT Pro	GAA Glu 50	GGC Gly	AAG Lys	AGT Ser	GTC Val	AAA Lys 55	GTT Val	TCG Ser	AAC Asn	TAC Tyr	CAC His 60	GTA Val	ATC Ile	270
CAC His	GGC Gly	AAG Lys 65	GAA Glu	TAC Tyr	TTT Phe	ATT Ile	GAA Glu 70	GGA Gly	ACT Thr	GCC Ala	TAC Tyr	CCA Pro 75	GTT Val	GGT Gly	315
GAC Asp	TCC Ser	AAG Lys 80	ATT Ile	GGA Gly	AAG Lys	ATC Ile	TAC Tyr 85	CAC His	AGC Ser	CTG Leu	ACT Thr	TAC Tyr 90	GGA Gly	GGT Gly	360
GTC Val	ACC Thr	AAG Lys 95	GAG Glu	AAC Asn	GTA Val	TTC Phe	AAC Asn 100	GTA Val	CTC Leu	TCC Ser	ACT Thr	GAC Asp 105	AAC Asn	AAG Lys	405
					TAC Tyr										450
GGA Gly	CAC His	CAA Gln 125	GAC Asp	TTC Phe	GTC Val	TGG Trp	GTG Val 130	CTC Leu	TCC Ser	AGA Arg	AGC Ser	ATG Met 135	GTC Val	CTT Leu	495
ACT Thr	GGT Gly	GAA Glu 140	GCC Ala	AAG Lys	ACC Thr	GCT Ala	GTC Val 145	GAG Glu	AAC Asn	TAC Tyr	CTT Leu	ATC Ile 150	GGC Gly	TCC Ser	540
CCA Pro	GTA Val	GTC Val 155	GAC Asp	TCC Ser	CAG Gln	AAA Lys	CTG Leu 160	GTA Val	TAC Tyr	AGT Ser	GAC Asp	TTC Phe 165	TCT Ser	GAA Glu	585
GCC Ala	GCC Ala	TGC Cys 170	AAG Lys	GTC Val	AAC Asn	AAT Asn	AGC Ser 175	AAC Asn	TGG Trp	TCT Ser	CAC His	CCG Pro 180	CAG Gln	TTC Phe	630
GAA Glu	AAA Lys	CCA Pro 185	GCT Ala	AGC Ser	CTG Leu	GCT Ala	GAA Glu 190	GCT Ala	AAA Lys	GTT Val	CTG Leu	GCT Ala 195	AAC Asn	CGT Arg	675
GAA Glu	CTG Leu	GAC Asp 200	AAA Lys	TAC Tyr	GGT Gly	GTT Val	TCC Ser 205	GAC Asp	TAC Tyr	TAC Tyr	AAA Lys	AAC Asn 210	CTC Leu	ATC Ile	720
AAC Asn	AAC Asn	GCT Ala 215	AAA Lys	ACC Thr	GTT Val	GAA Glu	GGT Gly 220	GTT Val	AAA Lys	GCT Ala	CTG Leu	ATC Ile 225	GAC Asp	GAA Glu	765

ATT CTC GCA GCA CTG CCG TAATAAGCTT
Ile Leu Ala Ala Leu Pro
230

793

INFORMATION FOR SEQ ID NO:10:

SEQUENCE CHARACTERISTICS:

LENGTH: 17 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:10:

GACGGTGCCT GTCCCGA 17

INFORMATION FOR SEQ ID NO:11:

SEQUENCE CHARACTERISTICS:

LENGTH: 17 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:11:

GACTACTGGG GAGCCGA 17

INFORMATION FOR SEQ ID NO:12:

SEQUENCE CHARACTERISTICS:

LENGTH: 522

TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: coding sequence of mutein DigA

FEATURE:

NAME/KEY: coding sequence

LOCATION: (1..522) OTHER INFORMATION:

/Product = "mutein without fusion parts"

SEQUENCE DESCRIPTION: SEQ ID NO:12:

GAC GTG TAC CAC GAC GGT GCC TGT CCC GAA GTC AAG CCA GTC GAC 45
Asp Val Tyr His Asp Gly Ala Cys Pro Glu Val Lys Pro Val Asp
1 5 10 15

AAC TTC GAC TGG TCC CAG TAC CAT GGT AAA TGG TGG GAA GTC GCC 90 Asn Phe Asp Trp Ser Gln Tyr His Gly Lys Trp Trp Glu Val Ala 20 25 30

AAA Lys	TAC	CCC	CAT His	CAC His 35	GAG Glu	CGG Arg	AAG Lys	TAC Tyr	GGA Gly 40	AAG Lys	TGC Cys	GGA Gly	TGG Trp	GCT Ala 45	135
GAG Glu	TAC Tyr	ACT Thr	CCT Pro	GAA Glu 50	GGC Gly	AAG Lys	AGT Ser	GTC Val	AAA Lys 55	GTT Val	TCG Ser	CGC Arg	TAC Tyr	TCT Ser 60	180
GTA Val	ATC Ile	CAC His	GGC Gly	AAG Lys 65	GAA Glu	TAC Tyr	TTT Phe	TCC Ser	GAA Glu 70	GGT Gly	ACC Thr	GCC Ala	TAC Tyr	CCA Pro 75	225
GTT Val	GGT Gly	GAC Asp	TCC Ser	AAG Lys 80	ATT Ile	GGA Gly	AAG Lys	ATC Ile	TAC Tyr 85	CAC His	AGC Ser	TAC Tyr	ACT Thr	ATT Ile 90	270
GGA Gly	GGT Gly	GTG Val	ACC Thr	CAG Gln 95	GAG Glu	GGT Gly	GTA Val	TTC Phe	AAC Asn 100	GTA Val	CTC Leu	TCC Ser	ACT Thr	GAC Asp 105	315
AAC Asn	AAG Lys	AAC Asn	TAC Tyr	ATC Ile 110	ATC Ile	GGA Gly	TAC Tyr	TTT Phe	TGC Cys 115	TCG Ser	TAC Tyr	GAC Asp	GAG Glu	GAC Asp 120	360
AAG Lys	AAG Lys	GGA Gly	CAC His	ATG Met 125	GAC Asp	TTG Leu	GTC Val	TGG Trp	GTG Val 130	CTC Leu	TCC Ser	AGA Arg	AGC Ser	ATG Met 135	405
GTC Val	CTT Leu	ACT Thr	GGT Gly	GAA Glu 140	GCC Ala	AAG Lys	ACC Thr	GCT Ala	GTC Val 145	GAG Glu	AAC Asn	TAC Tyr	CTT Leu	ATC Ile 150	450
GGC Gly	TCC Ser	CCA Pro	GTA Val	GTC Val 155	GAC Asp	TCC Ser	CAG Gln	AAA Lys	CTG Leu 160	GTA Val	TAC Tyr	AGT Ser	GAC Asp	TTC Phe 165	495
TCT Ser	GAA Glu	GCC Ala	GCC Ala	TGC Cys 170	AAG Lys	GTC Val	AAC Asn	AAT Asn							522

INFORMATION FOR SEQ ID NO:13:

SEQUENCE CHARACTERISTICS:

LENGTH: 76 bases TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

MOLECULE TYPE: synthetic oligodeoxynucleotide

SEQUENCE DESCRIPTION: SEQ ID NO:13:

CTGGTCCCAG TACCATGGTA AATGGTGGNN KGTCGCCNNK TACCCCNNKN 50 NKNNKNNKAA GTACGGAAAG TGCGGA 76

INFORMATION FOR SEQ ID NO:14:

SEQUENCE CHARACTERISTICS:

LENGTH: 1219 base pairs

TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: fragment of phasmid pBBP24

FEATURE:

NAME/KEY: signal peptide

LOCATION: (22..84)

FEATURE:

NAME/KEY: mature peptide LOCATION: (85..1209) OTHER INFORMATION:

/Product = "fusion protein composed of bilin-binding protein

Strep-tag II and fragment of phage envelope protein pIII, with interrupted reading frame"

/Codon = (sequence: "TAG", amino acid:Gln)

FEATURE:

NAME/KEY: coding sequence

LOCATION: (85..606) OTHER INFORMATION:

/Product = "mature bilin-binding protein with interrupted

reading frame"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (607..636)
OTHER INFORMATION:

/Product = "Strep-tag II affinity tag"

FEATURE:

NAME/KEY: coding sequence LOCATION: (637..639)

OTHER INFORMATION:

/Product = "amber stop codon

FEATURE:

NAME/KEY: coding sequence LOCATION: (640..1209)

OTHER INFORMATION:

/Product = "amino acids 217-406 of envelope protein pIII"

EQUENCE DESCRIPTION: SEQ ID NO:14:

TCTAGATAAC GAGGGCAAAA A ATG AAA AAG ACA GCT ATC GCG ATT

Met Lys Lys Thr Ala Ile Ala Ile

-21 -20 -15

GCA GTG GCA CTG GCT GGT TTC GCT ACC GTA GCG CAG GCC GAC GTG 90 Ala Val Ala Leu Ala Gly Phe Ala Thr Val Ala Gln Ala Asp Val -10 -5 -1 1

TAC Ty:	C CAG	C GAO S Ası	C GG7 p Gly 5	r GCC / Ala	TGT Cys	CCC Pro	GAA Glu 10	\Val	C AAG	CCA Pro	GT(GAC L Asp) Asr	TTC Phe	135	
GA(Ası	TG(G TCC Ser 20	GII	TAC Tyr	CAT His	GGT Gly	AAA Lys 25	Trp	TGG Trp	GAZ Glu	GTC Val	GCC Ala	Lys	TAC	180	
CCC Pro	AAC Asr	TCA Ser 35	. vai	GAG Glu	AAG Lys	TAC Tyr	GGA Gly 40	Asn	TAA	TGA	TGC	G GCT Ala 45	Glu	TAC Tyr	225	
1711	PIC	50	i GIĀ	, ràs	AGT Ser	Val	Lys 55	Val	Ser	Asn	Tyr	His 60	Val	Ile		
nis	GIY	г Був 65	GIU	туr	TTT Phe	Ile	Glu 70	Gly	Thr	Ala	Tyr	Pro 75	Val	Gly		
Asp	ser	80 rys	ITE	GIÀ	AAG Lys	Ile	Tyr 85	His	Ser	Leu	Thr	Tyr 90	Gly	Gly		
vai	TIII	95	GIU	ASn	GTA Val	Pne	100	Val	Leu	Ser	Thr	Asp 105	Asn	Lys		
ASII	ıyı	110	ile	GIĀ	TAC Tyr	TYY	Cys 115	Lys	Tyr	Asp	Glu	Asp 120	Lys	Lys		
GTĀ	nis	125	Asp	Pne	GTC Val	Trp	Val 130	Leu	Ser	Arg	Ser	Met 135	Val	Leu		
THE	GIÀ	140	Ala	Lys	ACC (Thr)	Ala	Val 145	Glu	Asn	Tyr	Leu	Ile 150	Gly	Ser		
Pro	vai	155	Asp	Ser	CAG A	Lys	Leu 160	Val	Tyr	Ser	Asp	Phe 165	Ser	Glu		
GCC Ala	GCC Ala	TGC Cys 170	AAG Lys	GTC Val	AAC A Asn A	Asn .	AGC Ser 175	AAC Asn	TGG Trp	TCT Ser	CAC His	CCG Pro 180	CAG Gln	TTC Phe	630	
GAA Glu	AAA Lys	TAG Gln 185	GCT Ala	GGC Gly	GGC (Hy S	TCT Ser 190	GGT Gly	GGT Gly	GGT Gly	TCT Ser	GGC Gly 195	GGC Gly	GGC Gly	675	
TCT Ser	Glu	GGT Gly 200	GGT Gly	GGC Gly	TCT (ilu (GGT (Gly (205	GGC Gly	GGT Gly	TCT Ser	GAG Glu	GGT Gly 210	GGC Gly	GGC Gly	720	

5	56.	r GA	G GG. u Gl; 21:	A GG(y Gl) 5	C GGT 7 Gly	TCC Ser	GG1 Gly	GG1 Gly 220	(GT)	C TC: y Sei	r GG1	TCC Ser	GGT Gly 225	' Asp	TTT Phe	765
	GAT Asp	TA!	r GAZ C Glu 230	A AAC 1 Lys	ATG Met	GCA Ala	AAC Asn	GCT Ala 235	L ASI	r AAC n Lys	GGG Gly	GCT Ala	ATO Met	Thr	GAA Glu	810
0	AAT Asn	GCC Ala	GAT ASI 245	GAA Glu	AAC Asn	GCG Ala	CTA Leu	CAG Gln 250	Ser	GAC Asp	GCT Ala	AAA Lys	GGC Gly 255	' Lys	CTT Leu	855
j	GAT Asp	TCT Ser	GTC Val 260	GCT Ala	ACT Thr	GAT Asp	TAC Tyr	GGT Gly 265	GCT Ala	GCT Ala	'ATC	GAT Asp	GGT Gly 270	TTC Phe	ATT Ile	900
1	GGT Gly	GAC Asp	GTT Val 275	TCC Ser	GGC Gly	CTT Leu	GCT Ala	AAT Asn 280	GGT Gly	`AAT 'Asn	GGT Gly	GCT Ala	ACT Thr 285	GGT Gly	GAT Asp	945
	TTT Phe	GCT Ala	GGC Gly 290	TCT Ser	AAT Asn	TCC Ser	CAA Gln	ATG Met 295	GCT Ala	CAA Gln	GTC Val	GGT Gly	GAC Asp 300	GGT Gly	GÁT Asp	990
	AAT Asn	TCA Ser	CCT Pro 305	TTA Leu	ATG Met	AAT Asn	AAT Asn	TTC Phe 310	CGT Arg	CAA Gln	TAT Tyr	TTA Leu	CCT Pro 315	TCC Ser	CTC Leu	1035
	CCT Pro	CAA Gln	TCG Ser 320	GTT Val	GAA Glu	TGT Cys	CGC Arg	CCT Pro 325	TTT Phe	GTC Val	TTT Phe	GGC Gly	GCT Ala 330	GGT Gly	AAA Lys	1080
	CCA Pro	TAT Tyr	GAA Glu 335	TTT Phe	TCT Ser	ATT Ile	GAT Asp	TGT Cys 340	GAC Asp	AAA Lys	ATA Ile	AAC Asn	TTA Leu 345	TTC Phe	CGT Arg	1125
	GGT Gly	GTC Val	TTT Phe 350	GCG Ala	TTT Phe	CTT Leu	Leu	TAT Tyr 355	GTT Val	GCC Ala	ACC Thr	Phe	ATG Met 360	TAT Tyr	GTA Val	1170
	TTT Phe	ser	ACG Thr 365	TTT Phe	GCT Ala A	AAC A Asn :	Ile :	CTG Leu 370	CGT Arg	AAT Asn	AAG Lys	Glu	TCT Ser 375			1209
•	TAAT.	AAGC	TT												;	1219

INFORMATION FOR SEQ ID NO:15:

SEQUENCE CHARACTERISTICS:

LENGTH: 522

TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: coding sequence of mutein DigA16

FEATURE:

NAME/KEY: coding sequence

LOCATION: (1..522) OTHER INFORMATION:

/Product = "mutein without fusion parts"

SEQUENCE DESCRIPTION: SEQ ID NO:15:

	CAC His							45
	TGG Trp							90
	GAT Asp						GCT Ala 45	135
	CCT Pro						TCT Ser 60	180
	GGC Gly						CCA Pro 75	225
 	 TCC Ser		 	 	 	 -	ATT Ile 90	270
	ACC Thr						GAC Asp 105	315
	TAC Tyr						GAC Asp 120	360
	CAC His						ATG Met 135	405
	GGT Gly						ATC Ile 150	450
	GTA Val						TTC Phe 165	495
	GCC Ala							522

INFORMATION FOR SEQ ID NO:16

SEQUENCE CHARACTERISTICS:

LENGTH: 1380 base pairs

TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: fragment of plasmid pBBP21

FEATURE:

NAME/KEY: signal peptide

LOCATION: (22..84)

FEATURE:

NAME/KEY: mature peptide

LOCATION: (85..636) OTHER INFORMATION:

/Product = "fusion protein composed of bilin-binding protein

Strep-tag II"

FEATURE:

NAME/KEY: signal peptide LOCATION: (658..717)

FEATURE:

NAME/KEY: mature peptide LOCATION: (718..1365) OTHER INFORMATION:

/Product = "DsbC protein"

SEQUENCE DESCRIPTION: SEQ ID NO:16:

TCTAGATAAC GAGGGCAAAA A ATG AAA AAG ACA GCT ATC GCG ATT

Met Lys Lys Thr Ala Ile Ala Ile

-21 -20 -15

GCA GTG GCA CTG GCT GGT TTC GCT ACC GTA GCG CAG GCC GAC GTG 90
Ala Val Ala Leu Ala Gly Phe Ala Thr Val Ala Gln Ala Asp Val
-10 -5 -1 1

TAC CAC GAC GGT GCC TGT CCC GAA GTC AAG CCA GTC GAC AAC TTC 135
Tyr His Asp Gly Ala Cys Pro Glu Val Lys Pro Val Asp Asn Phe
5 10 15

GAC TGG TCC CAG TAC CAT GGT AAA TGG TGG GAA GTC GCC AAA TAC 180 Asp Trp Ser Gln Tyr His Gly Lys Trp Trp Glu Val Ala Lys Tyr 20 25 30

CCC AAC TCA GTT GAG AAG TAC GGA AAG TGC GGA TGG GCT GAG TAC 225 Pro Asn Ser Val Glu Lys Tyr Gly Lys Cys Gly Trp Ala Glu Tyr 35 40 45

ACT CCT GAA GGC AAG AGT GTC AAA GTT TCG AAC TAC CAC GTA ATC 270 Thr Pro Glu Gly Lys Ser Val Lys Val Ser Asn Tyr His Val Ile 50 55 60

CAC His	GGC Gly	AAG Lys 65	GAA Glu	TAC Tyr	TTT Phe	ATT Ile	GAA Glu 70	GGA Gly	ACT Thr	GCC Ala	TAC Tyr	CCA Pro 75	GTT Val	GGT Gly	315
					AAG Lys										360
					GTA Val										405
					TAC Tyr										450
					GTC Val										495
					ACC Thr										540
					CAG Gln										585
					AAC Asn										630
	AAA Lys	TAA	raago	CTT (CGGG1	AAGA!	T T				GGT Gly			675	
					GCG Ala										720
					CAA Gln										765
					GCG Ala										810
					TTG Leu										855
					TAT Tyr										900

OTTO:	3.00	3300	7 7 C	a mc	cmc	mm z	220	C C C	mmc.	3 3 M	-	amm			0.45
Val	Thr	Asn	Lys 65	Met	Leu	Leu	Lys	Gln 70	Leu	Asn	Ala	Leu	GAA Glu 75	AAA Lys	945
			GTT Val 80											ACC Thr	990
			GAT Asp 95												1035
			GAC Asp 110												1080
			CGC Arg 125												1125
			TGG Trp 140												1170
			GGT Gly 155												1215
			CAT His 170												1260
			GTT Val 185												1305
			AAA Lys 200												1350
			GGT Gly 215		TAAT	TCGC	GT P	AGCT?	r.						1380

INFORMATION FOR SEQ ID NO:17:

SEQUENCE CHARACTERISTICS:

LENGTH: 2009 base pairs
TYPE: nucleic acid

STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: fragment of plasmid pBBP27

FEATURE:

NAME/KEY: signal peptide

LOCATION: (23..85)

FEATURE:

NAME/KEY: mature peptide LOCATION: (86..1999) OTHER INFORMATION:

FEATURE:

NAME/KEY: coding sequence LOCATION: (86..1435)
OTHER INFORMATION:

/Product = "linker peptide Pro-Pro-Ser-Ala"

FEATURE:

NAME/KEY: coding sequence LOCATION: (1448..1969) OTHER INFORMATION:

/Product = "DigA16 mutein"

FEATURE:

NAME/KEY: coding sequence LOCATION: (1970..1999) OTHER INFORMATION:

/Product = "Strep-tag II affinity tag"

SEQUENCE DESCRIPTION: SEQ ID NO:17:

TCTAGAACAT GGAGAAAATA AA GTG AAA CAA AGC ACT ATT GCA CTG

Val Lys Gln Ser Thr Ile Ala Leu

-21 -20 -15

GCA CTC TTA CCG TTA CTG TTT ACC CCT GTG ACA AAA GCC CGG ACA 91
Ala Leu Leu Pro Leu Leu Phe Thr Pro Val Thr Lys Ala Arg Thr
-10 -5 -1 1

CCA GAA ATG CCT GTT CTG GAA AAC CGG GCT GCT CAG GGC GAT ATT 136
Pro Glu Met Pro Val Leu Glu Asn Arg Ala Ala Gln Gly Asp Ile
5 10 15

ACT GCA CCC GGC GGT GCT CGC CGT TTA ACG GGT GAT CAG ACT GCC 181
Thr Ala Pro Gly Gly Ala Arg Arg Leu Thr Gly Asp Gln Thr Ala
20 25 30

GCT CTG CGT GAT TCT CTT AGC GAT AAA CCT GCA AAA AAT ATT ATT 226 Ala Leu Arg Asp Ser Leu Ser Asp Lys Pro Ala Lys Asn Ile Ile 35 40 45

TTG CTG ATT GGC GAT GGG ATG GGG GAC TCG GAA ATT ACT GCC GCA 271 Leu Leu Ile Gly Asp Gly Met Gly Asp Ser Glu Ile Thr Ala Ala 50 55 60

			GAA Glu										316
			ACC Thr										361
			CCG Pro										406
			ACC Thr										451
			GAA Glu										496
			CTG Leu										541
			CCC Pro										586
			CCG Pro										631
			GGC Gly										676
			GAC Asp										721
Glu	Thr	Ala	ACC Thr	Ala	Gly	Glu	Trp	Gln	Gly	Lys	Thr		766
			GCG Ala										811
			GTG Val										856
			GAC Asp										901

AAA Lys	GCA Ala	ACG Thr 275	TAC Tyr	CAT His	GGC Gly	AAT Asn	ATC Ile 280	GAT Asp	AAG Lys	CCC Pro	GCA Ala	GTC Val 285	ACC Thr	TGT Cys	946
												CTG Leu 300		CAG Gln	991
												GAG Glu 315			1036
												CAG Gln 330			1081
												GAT Asp 345			1126
GAA Glu	GCC Ala	GTA Val 350	CAA Gln	CGG Arg	GCG Ala	CTG Leu	GAA Glu 355	TTC Phe	GCT Ala	AAA Lys	AAG Lys	GAG Glu 360	GGT Gly	AAC Asn	1171
												AGC Ser 375			1216
GTT Val	GCG Ala	CCG Pro 380	GAT Asp	ACC Thr	AAA Lys	GCT Ala	CCG Pro 385	GGC Gly	CTC Leu	ACC Thr	CAG Gln	GCG Ala 390	CTA Leu	AAT Asn	1261
ACC Thr	AAA Lys	GAT Asp 395	GGC Gly	GCA Ala	GTG Val	ATG Met	GTG Val 400	ATG Met	AGT Ser	TAC Tyr	GGG Gly	AAC Asn 405	TCC Ser	GAA Glu	1306
												ATT Ile 420			1351
												GAC Asp 435			1396
GAT Asp	CTC Leu	TTC Phe 440	TAC Tyr	ACC Thr	ATG Met	AAA Lys	GCC Ala 445	GCT Ala	CTG Leu	GGG Gly	CTG Leu	AAA Lys 450	CCG Pro	CCT Pro	1441
AGC Ser	GCT Ala	GAC Asp 455	GTG Val	TAC Tyr	CAC His	GAC Asp	GGT Gly 460	GCC Ala	TGT Cys	CCC Pro	GAA Glu	GTC Val 465	AAG Lys	CCA Pro	1486
GTC Val	GAC Asp	AAC Asn 470	TTC Phe	GAC Asp	TGG Trp	TCC Ser	CAG Gln 475	TAC Tyr	CAT His	GGT Gly	AAA Lys	TGG Trp 480	Trp	CAG Gln	1531

				CCC Pro											1576
TGG Trp	GCT Ala	GAG Glu 500	TAC Tyr	ACT Thr	CCT Pro	GAA Glu	GGC Gly 505	AAG Lys	AGT Ser	GTC Val	AAA Lys	GTT Val 510	TCG Ser	CGC Arg	1621
TAC Tyr	TCT Ser	GTA Val 515	ATC Ile	CAC His	GGC Gly	AAG Lys	GAA Glu 520	TAC Tyr	TTT Phe	TCC Ser	GAA Glu	GGT Gly 525	ACC Thr	GCC Ala	1666
				GAC Asp											1711
				GTG Val											1756
				AAC Asn											1801
				GGA Gly											1846
AGC Ser	ATG Met	GTC Val 590	CTT Leu	ACT Thr	GGT Gly	GAA Glu	GCC Ala 595	AAG Lys	ACC Thr	GCT Ala	GTC Val	GAG Glu 600	AAC Asn	TAC Tyr	1891
				CCA Pro											1936
				GCC Ala											1981
				GAA Glu		TAATAAGCTT									2009

INFORMATION FOR SEQ ID NO:18:

SEQUENCE CHARACTERISTICS:

LENGTH: 2005 base pairs

TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear

MOLECULE TYPE: fragment of plasmid pBBP29

FEATURE:

NAME/KEY: signal peptide

LOCATION: (22..84)

FEATURE:

NAME/KEY: mature peptide LOCATION: (85..1998) OTHER INFORMATION:

/Product = "fusion protein composed of DigA16 mutein, Strep

tag II, linker peptide Gly(5) and alkaline

phosphatase"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (85..606) OTHER INFORMATION:

/Product = "DigAl6 mutein"

FEATURE:

NAME/KEY: coding sequence LOCATION: (607..636) OTHER INFORMATION:

/Product = "Strep-tag II affinity tag"

FEATURE:

NAME/KEY: coding sequence LOCATION: (637..651)

OTHER INFORMATION:

/Product = "linker peptide Gly-Gly-Gly-Gly"

FEATURE:

NAME/KEY: coding sequence

LOCATION: (652..1998)
OTHER INFORMATION:

/Product = "alkaline phosphatase without signal sequence and

N-terminal Arg"

SEQUENCE DESCRIPTION: SEQ ID NO:18:

TCTAGATAAC GAGGGCAAAA A ATG AAA AAG ACA GCT ATC GCG ATT

Met Lys Lys Thr Ala Ile Ala Ile

-21 -20 -15

GCA GTG GCA CTG GCT GGT TTC GCT ACC GTA GCG CAG GCC GAC GTG 90
Ala Val Ala Leu Ala Gly Phe Ala Thr Val Ala Gln Ala Asp Val
-10 -5 -1 1

TAC CAC GAC GGT GCC TGT CCC GAA GTC AAG CCA GTC GAC AAC TTC 135 Tyr His Asp Gly Ala Cys Pro Glu Val Lys Pro Val Asp Asn Phe 5 10 15

GAC TGG TCC CAG TAC CAT GGT AAA TGG TGG CAG GTC GCC GCG TAC 180 Asp Trp Ser Gln Tyr His Gly Lys Trp Trp Gln Val Ala Ala Tyr 20 25 30

CCC GAT CAT ATT ACG AAG TAC GGA AAG TGC GGA TGG GCT GAG TAC 225
Pro Asp His Ile Thr Lys Tyr Gly Lys Cys Gly Trp Ala Glu Tyr
35 40 45

ACT CCT GAA GGC AAG AGT GTC AAA GTT TCG CGC TAC TCT GTA ATC 270 Thr Pro Glu Gly Lys Ser Val Lys Val Ser Arg Tyr Ser Val Ile
50 55 60

CAC His	GGC Gly	AAG Lys 65	GAA Glu	TAC Tyr	TTT Phe	TCC Ser	GAA Glu 70	GGT Gly	ACC Thr	GCC Ala	TAC Tyr	CCA Pro 75	GTT Val	GGT Gly	315
GAC Asp	TCC Ser	AAG Lys 80	ATT Ile	GGA Gly	AAG Lys	ATC Ile	TAC Tyr 85	CAC His	AGC Ser	TAC Tyr	ACT Thr	ATT Ile 90	GGA Gly	GGT Gly	360
GTG Val	ACC Thr	CAG Gln 95	GAG Glu	GGT Gly	GTA Val	TTC Phe	AAC Asn 100	GTA Val	CTC Leu	TCC Ser	ACT Thr	GAC Asp 105	AAC Asn	AAG Lys	405
					TAC Tyr										450
GGA Gly	CAC His	ATG Met 125	GAC Asp	TTG Leu	GTC Val	TGG Trp	GTG Val 130	CTC Leu	TCC Ser	AGA Arg	AGC Ser	ATG Met 135	GTC Val	CTT Leu	495
ACT Thr	GGT Gly	GAA Glu 140	GCC Ala	AAG Lys	ACC Thr	GCT Ala	GTC Val 145	GAG Glu	AAC Asn	TAC Tyr	CTT Leu	ATC Ile 150	Gly	TCC Ser	540
					CAG Gln										585
					AAC Asn										630
					GGT Gly										675
AAC Asn	CGG Arg	GCT Ala 200	GCT Ala	CAG Gln	GGC Gly	GAT Asp	ATT Ile 205	ACT Thr	GCA Ala	CCC Pro	GGC Gly	GGT Gly 210	GCT Ala	CGC Arg	720
CGT Arg	TTA Leu	Thr	Gly	Asp	CAG Gln	Thr	Ala	Ala	Leu	Arg	Asp	Ser	Leu	AGC Ser	765
GAT Asp	AAA Lys	CCT Pro 230	GCA Ala	AAA Lys	AAT Asn	ATT Ile	ATT Ile 235	TTG Leu	CTG Leu	ATT Ile	Gly	GAT Asp 240	GGG Gly	ATG Met	810
					ACT Thr										855
					GGT Gly										900

TAC Tyr	ACT Thr	CAC His 275	TAT Tyr	GCG Ala	CTG Leu	AAT Asn	AAA Lys 280	AAA Lys	ACC Thr	GGC Gly	AAA Lys	CCG Pro 285	GAC Asp	TAC Tyr	945
GTC Val	ACC Thr	GAC Asp 290	TCG Ser	GCT Ala	GCA Ala	TCA Ser	GCA Ala 295	ACC Thr	GCC Ala	TGG Trp	TCA Ser	ACC Thr 300	GGT Gly	GTC Val	990
AAA Lys	ACC Thr	TAT Tyr 305	AAC Asn	GGC Gly	GCG Ala	CTG Leu	GGC Gly 310	GTC Val	GAT Asp	ATT Ile	CAC His	GAA Glu 315	AAA Lys	GAT Asp	1035
CAC His	CCA Pro	ACG Thr 320	ATT Ile	CTG Leu	GAA Glu	ATG Met	GCA Ala 325	AAA Lys	GCC Ala	GCA Ala	GGT Gly	CTG Leu 330	GCG Ala	ACC Thr	1080
GGT Gly	AAC Asn	GTT Val 335	TCT Ser	ACC Thr	GCA Ala	GAG Glu	TTG Leu 340	CAG Gln	GAT Asp	GCC Ala	ACG Thr	CCC Pro 345	GCT Ala	GCG Ala	1125
CTG Leu	GTG Val	GCA Ala 350	CAT His	GTG Val	ACC Thr	TCG Ser	CGC Arg 355	AAA Lys	TGC Cys	TAC Tyr	GGT Gly	CCG Pro 360	AGC Ser	GCG Ala	1170
ACC Thr	AGT Ser	GAA Glu 365	AAA Lys	TGT Cys	CCG Pro	GGT Gly	AAC Asn 370	GCT Ala	CTG Leu	GAA Glu	AAA Lys	GGC Gly 375	GGA Gly	AAA Lys	1215
GGA Gly	TCG Ser	ATT Ile 380	ACC Thr	GAA Glu	CAG Gln	CTG Leu	CTT Leu 385	AAC Asn	GCT Ala	CGT Arg	GCC Ala	GAC Asp 390	GTT Val	ACG Thr	1260
CTT Leu	GGC Gly	GGC Gly 395	GGC Gly	GCA Ala	AAA Lys	ACC Thr	TTT Phe 400	GCT Ala	GAA Glu	ACG Thr	GCA Ala	ACC Thr 405	GCT Ala	GGT Gly	1305
GAA Glu	TGG Trp	CAG Gln 410	GGA Gly	AAA Lys	ACG Thr	CTG Leu	CGT Arg 415	GAA Glu	CAG Gln	GCA Ala	CAG Gln	GCG Ala 420	CGT Arg	GGT Gly	1350
				AGC Ser											1395
				AAA Lys											1440
				TGG Trp											1485
				GCA Ala											1530

GAC Asp	AGT Ser	GTA Val 485	CCA Pro	ACC Thr	CTG Leu	GCG Ala	CAG Gln 490	ATG Met	ACC Thr	GAC Asp	AAA Lys	GCC Ala 495	ATT Ile	GAA Glu	1575
TTG Leu	TTG Leu	AGT Ser 500	AAA Lys	AAT Asn	GAG Glu	AAA Lys	GGC Gly 505	TTT Phe	TTC Phe	CTG Leu	CAA Gln	GTT Val 510	GAA Glu	GGT Gly	1620
GCG Ala	TCA Ser	ATC Ile 515	GAT Asp	AAA Lys	CAG Gln	GAT Asp	CAT His 520	GCT Ala	GCG Ala	AAT Asn	CCT Pro	TGT Cys 525	GGG Gly	CAA Gln	1665
ATT Ile	GGC Gly	GAG Glu 530	ACG Thr	GTC Val	GAT Asp	CTC Leu	GAT Asp 535	GAA Glu	GCC Ala	GTA Val	CAA Gln	CGG Arg 540	GCG Ala	CTG Leu	1710
					GAG Glu										1755
GAT Asp	CAC His	GCC Ala 560	CAC His	GCC Ala	AGC Ser	CAG Gln	ATT Ile 565	GTT Val	GCG Ala	CCG Pro	GAT Asp	ACC Thr 570	AAA Lys	GCT Ala	1800
CCG Pro	GGC Gly	CTC Leu 575	ACC Thr	CAG Gln	GCG Ala	CTA Leu	AAT Asn 580	ACC Thr	AAA Lys	GAT Asp	GGC Gly	GCA Ala 585	GTG Val	ATG Met	1845
GTG Val	ATG Met	AGT Ser 590	TAC Tyr	GGG Gly	AAC Asn	TCC Ser	GAA Glu 595	GAG Glu	GAT Asp	TCA Ser	CAA Gln	GAA Glu 600	CAT His	ACC Thr	1890
GGC Gly	AGT Ser	CAG Gln 605	TTG Leu	CGT Arg	ATT Ile	GCG Ala	GCG Ala 610	TAT Tyr	GGC Gly	CCG Pro	CAT His	GCC Ala 615	GCC Ala	AAT Asn	1935
GTT Val	GTT Val	GGA Gly 620	CTG Leu	ACC Thr	GAC Asp	CAG Gln	ACC Thr 625	GAT Asp	CTC Leu	TTC Phe	TAC Tyr	ACC Thr 630	ATG Met	AAA Lys	1980
		CTG Leu 635			AAA Lys	TAAG	CTT						`		2005

Patent Claims

- 1. A polypeptide selected from muteins of the bilinbinding protein, characterized in that it
- 5 (a) is able to bind digoxigenin or digoxigenin conjugates,
 - (b) does not bind ouabain, testosterone and 4-aminofluorescein and
- (c) has an amino acid substitution at at least one of the sequence positions 28, 31, 34, 35, 36, 37, 58, 60, 69, 88, 90, 95, 97, 114, 116, 125 and 127 of the bilin-binding protein.
- The polypeptide as claimed in claim 1,
 characterized in that the dissociation constant of the complex with digoxigenin is 100 nM or less.
- 3. The polypeptide as claimed in claim 1 or 2, characterized in that it carries, when compared to the bilin-binding protein, at least one of the amino acid substitutions selected from Glu(28)->Gln, Lys(31)->Ala, Asn(34)->Asp, Ser(35)->His, Val(36)->Ile, Glu(37)->Thr, Asn(58)->Arg, His(60)->Ser, Ile(69)->Ser, Leu(88)->Tyr, Tyr(90)->Ile, Lys(95)->Gln, Asn(97)->Gly,
- 25 Tyr(114)->Phe, Lys(116)->Ser, Gln(125)->Met and Phe(127)->Leu.
- 4. The polypeptide as claimed in one or more of claims 1 to 3, characterized in that it carries at least one label group, selected from enzymatic label, radioactive label, fluorescent label, chromophoric label, (bio)luminescent label or label containing haptens, biotin, metal complexes, metals or colloidal gold.

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5. A fusion protein of polypeptides as claimed in one or more of claims 1 to 4, characterized in that an enzyme, another protein or a protein domain, a signal

sequence and/or an affinity peptide is fused to the amino terminus of the polypeptide in an operable manner.

- 5 6. A fusion protein of polypeptides as claimed in one or more of claims 1 to 5, characterized in that an enzyme, another protein or a protein domain, a targeting sequence and/or an affinity peptide is fused to the carboxy terminus of the polypeptide in an operable manner.
- 7. A nucleic acid, characterized in that it comprises a sequence coding for a mutein or a fusion protein of a mutein of the bilin-binding protein as claimed in one or more of claims 1 to 6.
- 8. A method for preparing a mutein or a fusion protein of a mutein of the bilin-binding protein as claimed in one or more of claims 1 to 6, characterized in that the nucleic acid coding for the mutein or the fusion protein of a mutein of the bilin-binding protein is expressed in a bacterial or eukaryotic host cell and the polypeptide is obtained from the cell or the culture supernatant.

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- The use of a mutein or a fusion protein of a mutein of the bilin-binding protein as claimed in one or more of claims 1 to 8 for binding, detecting, determining, immobilizing or removing digoxigenin or conjugates of digoxigenin with proteins, nucleic acids, carbohydrates, other biological or synthetic macromolecules or low molecular weight chemical compounds.
- 35 10. A method for detecting the digoxigenin group, wherein a mutein of the bilin-binding protein or a fusion protein of a mutein of the bilin-binding protein as claimed in one or more of claims 1 to 8 is contacted with digoxigenin or

with conjugates of digoxigenin under conditions suitable for effecting binding of the mutein to the digoxigenin group, and the mutein or the fusion protein of the mutein is determined.

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⁴ pages(s) of drawings attached

Number: DE 199 26 068 C1
Int. Cl. 7: C 07 K 14/435

Published on: January 11, 2001

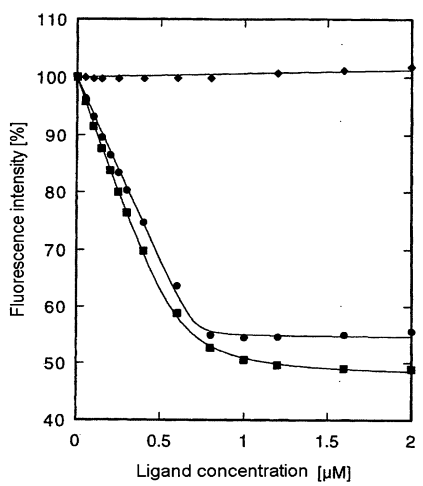
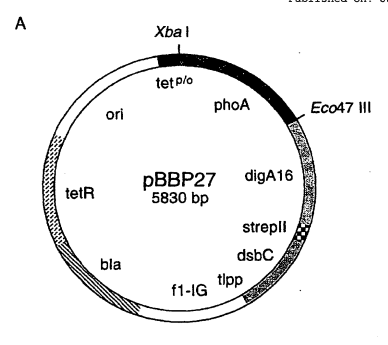


Figure 1

Number: DE 199 26 068 C1
Int. Cl. 7: C 07 K 14/435
Published on: January 11, 2001



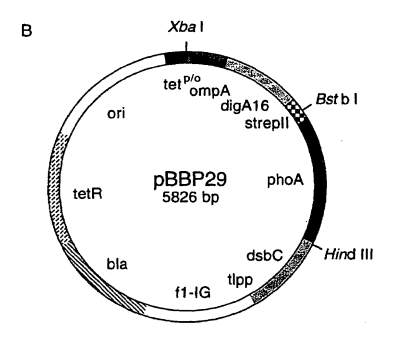


Figure 2

Number: DE 199 26 068 C1
Int. Cl. 7: C 07 K 14/435
Published on: January 11, 2001

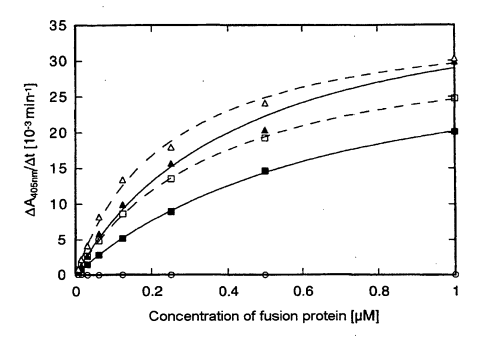


Figure 3

Number: DE 199 26 068 C1
Int. Cl. 7: C 07 K 14/435
Published on: January 11, 2001



Figure 4